

FORM PTO-1390 (Modified)
(Rev. 11-98)

DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE

ATTORNEY'S DOCKET NUMBER

TRANSMITTAL LETTER TO THE UNITED STATES

294-86 PCT/US

DESIGNATED/ELECTED OFFICE (DO/EO/US)

U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR

CONCERNING A FILING UNDER 35 U.S.C. 371

09/ 674752

INTERNATIONAL APPLICATION NO.

INTERNATIONAL FILING DATE

PRIORITY DATE CLAIMED

PCT/NL99/00285

7 May 1999 (07.05.99)

8 May 1998 (08.05.98)

TITLE OF INVENTION

METHOD FOR DIAGNOSIS AND TREATMENT OF HAEMOPHILIA A PATIENTS WITH AN INHIBITOR

APPLICANT(S) FOR DO/EO/US

Johannes Jacobus Voorberg, et al.

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

1. ☒ This is a **FIRST** submission of items concerning a filing under 35 U.S.C. 371.
2. ☐ This is a **SECOND** or **SUBSEQUENT** submission of items concerning a filing under 35 U.S.C. 371.
3. ☒ This is an express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1).
4. ☒ A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.
5. ☒ A copy of the International Application as filed (35 U.S.C. 371 (c) (2))
 - a. ☐ is transmitted herewith (required only if not transmitted by the International Bureau).
 - b. ☒ has been transmitted by the International Bureau.
 - c. ☐ is not required, as the application was filed in the United States Receiving Office (RO/US).
6. ☐ A translation of the International Application into English (35 U.S.C. 371(c)(2)).
7. ☒ A copy of the International Search Report (PCT/ISA/210).
8. ☒ Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371 (c)(3))
 - a. ☐ are transmitted herewith (required only if not transmitted by the International Bureau).
 - b. ☒ have been transmitted by the International Bureau.
 - c. ☐ have not been made; however, the time limit for making such amendments has NOT expired.
 - d. ☐ have not been made and will not be made.
9. ☐ A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
10. ☐ An oath or declaration of the inventor(s) (35 U.S.C. 371 (c)(4)).
11. ☒ A copy of the International Preliminary Examination Report (PCT/IPEA/409).
12. ☐ A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371 (c)(5)).

Items 13 to 20 below concern document(s) or information included:

13. ☐ An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
14. ☐ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
15. ☐ A **FIRST** preliminary amendment.
16. ☐ A **SECOND** or **SUBSEQUENT** preliminary amendment.
17. ☐ A substitute specification.
18. ☐ A change of power of attorney and/or address letter.
19. ☒ Certificate of Mailing by Express Mail
20. ☒ Other items or information:

Copy of the international application as published on 18 November 1999 under publication number WO 99/58680

U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR 1.53) 09/ 674752	INTERNATIONAL APPLICATION NO. PCT/NL99/00285	ATTORNEY'S DOCKET NUMBER 294-86 PCT/US
---	--	--

21. The following fees are submitted..

BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)) :

- ☐ Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO **\$1,000.00**
- ☒ International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO **\$860.00**
- ☐ International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO **\$710.00**
- ☐ International preliminary examination fee paid to USPTO (37 CFR 1.482) but all claims did not satisfy provisions of PCT Article 33(1)-(4) **\$690.00**
- ☐ International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(1)-(4) **\$100.00**

ENTER APPROPRIATE BASIC FEE AMOUNT =

Surcharge of **\$130.00** for furnishing the oath or declaration later than ☐ 20 ☐ 30 months from the earliest claimed priority date (37 CFR 1.492 (e)).

CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE	
Total claims	23 - 20 =	3	x \$18.00	\$54.00
Independent claims	4 - 3 =	1	x \$80.00	\$80.00
Multiple Dependent Claims (check if applicable) <input checked="" type="checkbox"/>				\$270.00
TOTAL OF ABOVE CALCULATIONS =				\$1,264.00
Reduction of 1/2 for filing by small entity, if applicable. Verified Small Entity Statement must also be filed (Note 37 CFR 1.9, 1.27, 1.28) (check if applicable) <input type="checkbox"/>				\$0.00
SUBTOTAL =				\$1,264.00
Processing fee of \$130.00 for furnishing the English translation later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492 (f)).				\$0.00
TOTAL NATIONAL FEE =				\$1,264.00
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31) (check if applicable) <input type="checkbox"/>				\$0.00
TOTAL FEES ENCLOSED =				\$1,264.00
				Amount to be refunded
				charged

- ☒ A check in the amount of **\$1,264.00** to cover the above fees is enclosed.
- ☐ Please charge my Deposit Account No. _____ in the amount of _____ to cover the above fees.
A duplicate copy of this sheet is enclosed.
- ☒ The Commissioner is hereby authorized to charge any fees which may be required, or credit any overpayment to Deposit Account No. **08-2461** A duplicate copy of this sheet is enclosed.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

SEND ALL CORRESPONDENCE TO:

Ronald J. Baron, Esq.
Hoffmann & Baron, LLP
6900 Jericho Turnpike
Syosset, New York 11791
United States of America

Phone (516) 822-3550
Facsimile (516) 822-3582

SIGNATURE

Ronald J. Baron

NAME

29,281

REGISTRATION NUMBER

November 6, 2000

DATE

PCT/NOV 2000

METHODS FOR DIAGNOSIS AND TREATMENT OF HAEMOPHILIA A PATIENTS WITH AN INHIBITOR

FIELD OF THE INVENTION

This invention is in the fields of diagnosis and medical treatment. More in particular, the invention provides means and methods for diagnosing the presence of inhibitory antibodies directed against factor VIII in the blood of human individuals, and provides means, pharmaceutical compositions and methods for treating human individuals in which such inhibitory antibodies occur.

BACKGROUND OF THE INVENTION

Haemophilia A is an X-linked bleeding disorder which is characterized by the functional absence of blood coagulation factor VIII. Depending on the residual factor VIII activity in the plasma of the patient, haemophilia A can be classified as severe (factor VIII < 1%), moderate (factor VIII 1-5%) or mild (>5%). Bleeding episodes in patients with haemophilia A can be effectively controlled by intravenous administration of purified factor VIII concentrates. These factor VIII-concentrates may be derived from pools of human plasma. Alternatively, recombinant factor VIII produced by genetically engineered eukaryotic cells may be used as a starting material for the preparation of factor VIII concentrates.

A serious complication of current haemophilia A treatment constitutes the development of neutralizing antibodies directed against factor VIII. These antibodies, commonly termed factor VIII inhibitors, arise in approximately 25% of the patients with severe haemophilia A, usually after 5-20 exposure-days (Ehrenforth et al. 1992, Lancet 339: 594-598). In patients with moderate and mild haemophilia A, anti-factor VIII antibodies occur less frequently and this is most likely due to induction of tolerance by endogenous factor VIII present in the plasma of this group of patients (McMillan et al. 1988, Blood 71: 344-348). Antibodies to factor VIII may develop with low frequency in healthy individuals.

Diagnosis of factor VIII inhibitors is commonly performed using the so-called Bethesda assay (Kasper et al. 1975, Thromb. Diath. Haemorrh. 34: 869-

872). In this assay equal amounts of normal plasma and dilutions of inhibitor plasma are incubated for two hours at 37°C. Next, residual factor VIII activity is determined and compared to control incubation in which normal plasma is incubated with 0.1 M imidazole for 2 hours at 37°C. The amount of inhibitor is expressed in Bethesda units; one Bethesda unit corresponds to the amount of inhibitor that is capable of reducing the activity of factor VIII in normal plasma with 50%. A recent study has proposed several adaptations to the original assay system which serve to improve the stability of factor VIII during the assay (Verbruggen et al. 1995, *Thromb. Haemostas.* 73: 247-251). This so-called "Nijmegen modification" of the Bethesda assay is particularly useful for the detection of low titre factor VIII inhibitors. It should be noted that the Bethesda assay does not provide information on the epitopes of factor VIII inhibitory antibodies.

The occurrence of factor VIII-inhibiting antibodies renders factor VIII replacement therapy inadequate. Several treatment options are available to the clinician. Low titre inhibitors (up to 5-10 BU/ml) are usually treated with infusion of high doses of factor VIII. A subset of factor VIII inhibitors does not cross react with porcine factor VIII. Porcine factor VIII has been used for management of patients with an inhibitor. Administration of porcine factor VIII may present with side effects. After multiple treatment 30-50% of the patients develop antibodies that neutralize the activity of the administered porcine factor VIII.

An alternative treatment of patients with factor VIII inhibitor constitutes the use of factor VIII bypassing agents. Activated prothrombin concentrates complexes (APCC) have been used to bypass the activity of factor VIII. APCC has been used successfully to control bleeding episodes in a large number of patients with an inhibitor. However, treatment is not effective in all cases and an anamnestic rise in the titre of the inhibitor following administration of APCC (most likely due to trace amounts of factor VIII in the preparation) has been reported in a number of patients. In the last 5 years recombinant factor VIIa has become available as a new factor VIII bypassing agent for the treatment of patients with an inhibitor (Lusher et al. 1996, *Haemostasis* 26 (suppl. 1): 124-130). Recombinant factor VIIa has been successfully used to control the bleeding episodes in patients with an inhibitor. Treatment by this agent is however limited by the short half-life of this compound in the circulation which requires multiple infusions at relatively short time intervals. APC-resistant factor V has recently

been suggested as an alternative means to bypass the biological activity of factor VIII inhibitors (WO 95/29259). The agents described above do not act directly on factor VIII inhibitors but merely serve to bypass factor VIII by infusion of large amounts of clotting factor concentrates with increased procoagulant activity.

Other methods of inhibitor neutralization have been proposed but their effectiveness has not been convincingly shown. Immunoglobulin preparations derived from plasma of healthy donors has been proposed as an active suppressor of factor VIII inhibitors (Sultan et al. 1984, Lancet 333, 765-768). Despite success in patients with acquired haemophilia A and high titre inhibitors, immunoglobulin preparations are not applied universally for treatment of patients with an inhibitor. The beneficial effects of immunoglobulin preparations in these patients have been attributed to the presence of anti-idiotypic antibodies that neutralize the activity of factor VIII inhibitors. Indeed in some patients the decline in the level of factor VIII inhibitors coincided with the appearance of anti-idiotypic antibodies (Sultan et al. 1987, Proc. Natl. Acad. Sci. USA 84: 828-831). Control of factor VIII inhibitors by anti-idiotypic antibodies in both haemophilia A patients and healthy individuals has been strongly advocated by some investigators (Gilles et al. 1996, J. Clin. Inv. 97: 1382-1388). The same group has proposed that infusion of antigen-antibody complexes in patients with inhibitors may accelerate a decline in anti-factor VIII antibodies in patients with an inhibitor (USP 5,543,145). It has been proposed that this decline is mediated by an increase in the number of anti-idiotypic antibodies which are induced by the infused antigen-antibody complexes. The factor VIII specific antibody used in this treatment protocol is derived from plasma of patients with an inhibitor. Obviously, this presents a heterogeneous mixture of antibodies and no details with respect to the epitope specificity of these antibodies are available. Also the primary structure of the antibodies in these antigen-antibody preparations has not been disclosed.

SUMMARY OF THE INVENTION

This invention relates to methods for diagnosis and treatment of inhibitory antibodies directed against factor VIII. Methods are disclosed that show how to arrive at nucleotide and amino acid sequences that encode factor VIII specific antibodies. This invention discloses diagnostic tests that allow for detection of nucleotide and amino acid sequences that encode factor VIII specific antibodies

within a heterogeneous mixture of antibody-encoding nucleotide or amino acid sequences. This invention further discloses how to use recombinant antibody fragments which bind specifically to factor VIII as novel therapeutic agents for the treatment of patients with factor VIII inhibitors.

5 The invention provides a polynucleotide in substantially isolated form, comprising a contiguous nucleotide sequence (a) coding for a human antibody with factor VIII specificity, or (b) complementary to a nucleotide sequence coding for a human antibody with factor VIII specificity, or (c) capable of selectively hybridizing under stringent conditions to nucleotide sequence (a) or (b).

10 Preferably, the contiguous nucleotide sequence is at least 8, preferably at least 10 nucleotides.

In a preferred embodiment, the invention provides a probe or primer which comprises a polynucleotide as defined herein, optionally further comprising a detectable label, such as a radioactive atom or group, an enzyme, a fluorescent or
15 luminescent group, a dye or biotin.

The invention also provides an assay kit for detecting nucleic acid coding for a human antibody with factor VIII specificity, comprising a probe or primer as defined herein in a suitable container.

Furthermore, the invention provides a nucleic acid amplification and
20 detection kit for detecting nucleic acid coding for a human antibody with factor VIII specificity, comprising a pair of primers as defined herein capable of priming the synthesis of cDNA, and optionally further comprising a probe as defined herein capable of selectively hybridizing to (the complement of) a region of the nucleic acid to be detected between and not including the sequences from
25 which the primers are derived.

The invention provides a method for assaying a sample for the presence or absence of nucleic acid coding for a human antibody with factor VIII specificity, comprising contacting the sample with a probe as defined herein under conditions that allow the selective hybridization of said probe to the (complement of
30 the) nucleic acid to be detected in the sample, and determining whether polynucleotide duplexes comprising said probe are formed.

The invention also provides a method for assaying a sample for the presence or absence of nucleic acid coding for a human antibody with factor VIII specificity, comprising subjecting nucleic acid present in the sample to a nucleic
35 acid amplification process using a pair of primers as defined herein capable of priming the synthesis of cDNA, contacting the nucleic acid resulting from the

amplification process with a probe as defined herein under conditions that allow the selective hybridization of said probe to the (complement of the) nucleic acid to be detected in the sample, and determining whether polynucleotide duplexes comprising said probe are formed.

5 Furthermore, the invention provides a method of producing a recombinant polypeptide, comprising providing a polynucleotide coding for said polypeptide, preparing a recombinant vector containing said polynucleotide operably linked to a control sequence capable of providing for the expression of the polynucleotide by a host cell, transforming a host cell with said recombinant vector, growing
10 said host cell under conditions that provide for the expression of the polynucleotide and optionally isolating the thus produced polypeptide, wherein said polynucleotide codes for a human antibody with factor VIII specificity, or a fragment or derivative thereof capable of specific binding to factor VIII.

According to another aspect, the invention provides a polypeptide in
15 substantially isolated form, comprising a contiguous amino acid sequence corresponding to or mimicking a fragment or derivative of a human antibody with factor VIII specificity capable of specific binding to factor VIII. In a preferred embodiment of the invention, the contiguous amino acid sequence is capable of reducing the activity of factor VIII inhibiting antibodies.

20 Preferably, the fragment is (part of) a variable region of the heavy chain or light chain of said antibody, and the derivative is preferably a single chain Fv fragment of said antibody.

The invention furthermore provides an antibody in substantially isolated form, comprising a recombinant human antibody with factor VIII specificity or
25 an anti-idiotypic antibody directed against a human antibody with factor VIII specificity.

The invention furthermore provides a pharmaceutical composition for the treatment of factor VIII inhibition in a human individual, comprising a polypeptide as defined herein or an antibody as defined herein, together with a
30 pharmaceutically acceptable carrier. Optionally, the composition further contains factor VIII, or a substitute of factor VIII.

The invention also provides a method of treatment of factor VIII inhibition in a human individual comprising administering (an effective amount to reduce or prevent said factor VIII inhibition of) a polypeptide as defined herein or an
35 antibody as defined herein, optionally together with factor VIII or a substitute of factor VIII.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the light chain specificity of 12 clones obtained after 4 rounds of panning of the IgG4-specific library described in Example 2. Phage expressing recombinant antibodies were incubated on microtiter wells which contained factor VIII light chain (black bars; +80K). To correct for background binding phage were also incubated on microtiter wells that did not contain factor VIII light chain (grey bars; -80K). On the Y-axis the OD(450-540 nm) is depicted. Two clones (c) express antibody fragments that do not bind specifically to the factor VIII light chain.

Figure 2 shows the light chain specificity of 12 clones randomly chosen after the first round of panning of the IgG4-specific library described in Example 2. Clones 7, 8 and 10 express antibody fragments with factor VIII light chain specificity. The other clones do not specifically bind to the factor VIII light chain. Only background binding of the phage to the microtiter wells is observed (grey bars). Two clones (c) express antibody fragments that do not bind specifically to the factor VIII light chain.

Figure 3 shows the nucleotide sequence of clone EL14 and clone IT2. The nucleotide sequence of both clones is aligned with the nucleotide sequence of the germline sequences DP-10 (for EL14) and DP-14 (for IT2). The different regions of the variable part of the heavy chain are indicated in the following order: framework 1, CDR1, framework 2, CDR2, framework 3, CDR3 and framework 4. Homology of clones EL14 and IT2 with the germline sequences DP-10 and DP-14 is indicated by horizontal bars (-). Differences are indicated by the nucleotides that occur in the germline sequences DP-10 and DP-14. Note that both CDR3 and framework 4 are not derived from the germline sequences DP-10 and DP-14. Consequently, no homology is given for this part of the nucleotide sequence.

Figure 4A gives the amino acid sequence derived of the nucleotide sequence of clone EL14 and IT2. Deviations in the amino acid sequence of the germline segments DP-10 and DP-14 are indicated in the lower lines. Framework is abbreviated as "FR".

Figure 4B compares the amino acid sequence of three related clones that are derived from the germline segment DP-14. The amino acid sequences of clone IT2, clone EL5 and clone EL25 are compared to that of the germline segment DP-14. Deviations in amino acid sequence are indicated for each clone. Note that some amino acid substitutions are shared by the three different clones.

Figure 4C compares the amino acid sequences of the third variable loop (CDR3) of the heavy chain of clone EL14 and IT2. Homologous amino acid residues are indicated by vertical lines. Dots denote amino acids related in charge or hydrophobicity.

5 Figure 5 shows the specificity of binding of scFv-EL14 and scFv-IT2 to the factor VIII light chain as assessed by the murine monoclonal antibodies CLB-CAG A and CLB-CAG 117. ScFv-EL14 binds specifically to the factor VIII light chain when peroxidase labelled CLB-CAG A (80K/Apo) is used as an indicator antibody (hatched bars). Also scFv-IT2 binds to the factor VIII light chain under
10 these conditions (hatched bars). In contrast, when peroxidase labelled CLB-CAG 117 (80K/117po) is used as indicator antibody binding of scFv-EL14 and scFv-IT2 is strongly reduced (black bars). No binding is observed in the absence of factor VIII light chain (-/117po; -/Apo). Clone O4 does not bind to factor VIII under these experimental conditions. These experiments show that the epitope of
15 scFv-EL14 and scFv-IT2 overlaps with that of CLB-CAG 117. On the y-axis the absorbance OD (450-540 nm) is given. On the x-axis scFv-EL14, scFv-IT2 and scFv-O4 are given.

Figure 6 shows the binding of different dilutions of purified scFv-EL14 (open circles), scFv-IT2 (closed circles) and scFv-O4 (negative control). On the
20 x-axis the different concentrations of protein tested are indicated ($\mu\text{g/ml}$), on the y-axis the absorbance OD(450-540 nm) is given. Clone scFv-O4 does not bind to the factor VIII light chain at the protein concentration tested in this experiment. Both scFv-IT2 and scFv-EL14 bind to the factor VIII light chain. ScFv-EL14 binds with a higher affinity to the factor VIII light chain when compared to scFv-
25 IT2.

Figure 7A shows the neutralization of the inhibitory activity of the murine monoclonal antibody CLB-CAG 117 by scFv-EL14. Antibody CLB-CAG 117 was diluted till a value of 2 BU/ml which corresponds with a residual factor VIII activity of about 25%. Increasing amounts of scFv-EL14 were capable of
30 neutralizing the inhibitory activity of CLB-CAG 117 (closed circles). A concentration of 0.75 $\mu\text{g/ml}$ suffices to restore factor VIII activity to its original level. ScFv-EL14 did not affect the inhibitory activity of the murine monoclonal antibody CLB-CAG A (open circles). On the y-axis residual factor VIII activity is depicted. On the x-axis the amount of scFv added is given in $\mu\text{g/ml}$.

35 Figure 7B shows the neutralization of the inhibitory activity of CLB-CAG 117 by scFv-IT2 (closed circles). A concentration of 65 $\mu\text{g/ml}$ is needed to

restore factor VIII activity to its original level. ScFv-IT2 did not affect the inhibitory activity of CLB-CAG A (open bars). On the y-axis residual factor VIII activity is depicted. On the x-axis the amount of scFv added is given in $\mu\text{g/ml}$.

Figure 8A shows the epitope specificity of 15 clones obtained after four rounds of panning of the IgG4-specific library described in example 8. Panning was performed using factor VIII immobilized on immunotubes. Phage expressing recombinant antibodies were incubated on microtiter wells which contained factor VIII light chain (black bars) or A3-C1-domain (hatched bars). To correct for background binding, phage were also incubated on microtiter wells that did not contain factor VIII light chain or A3-C1 domain (white bars). On the y-axis the OD (450-540 nm) is depicted.

Figure 8B shows the epitope specificity of 15 clones obtained after four rounds of panning the IgG4 specific library described in example 8. Panning was performed using factor VIII light chain that had been immobilized in microtiter wells employing CLB-CAG 117. Phage expressing recombinant antibodies were incubated on microtiter wells which contained factor VIII light chain (black bars) or A3-C1-domain (hatched bars). To correct for background binding, phage were also incubated on microtiter wells that did not contain factor VIII light chain or A3-C1 domain (white bars). On the y-axis the OD (450-540 nm) is depicted.

Figure 9A shows the deduced amino acid sequence of recombinant antibody fragments specific for the A3-C1 domain. The amino acid sequence of germ line variable heavy chain gene segments DP15, DP31, DP49 and DP77 is given. Deviations in amino acid sequence from these germline gene segments are indicated for clone B38, B18, B35 and B04. Also the amino acid of the CDR3 and FR4 of the A3-C1 specific recombinant antibodies encoded by clone B38, B18, B35 and B04 is given.

Figures 9B-E give the nucleotide and amino acid sequence of the variable heavy chain domain of clone B38, B18, B35 and B04.

Figure 10A shows the epitope specificity of 20 clones obtained after four rounds of panning of the IgG4-specific library described in Example 9. Panning was performed using factor VIII heavy chain (90K+) that had been immobilized in microtiter wells employing CLB-CAG 9. Phage expressing recombinant antibodies were incubated on microtiter wells which contained factor VIII heavy chain that was immobilized using the anti-heavy chain monoclonal antibody ESH5 (black bars). To correct for background binding,

phage were also incubated on microtiter wells that did not contain factor VIII heavy chain (white bars). On the y-axis the OD (450-540 nm) is depicted.

Figure 10B shows the epitope specificity of 20 clones obtained after four rounds of panning of the IgG4-specific library described in Example 9. Panning was performed using factor VIII heavy chain (90K+) that had been immobilized in immunotubes. Phage expressing recombinant antibodies were incubated on microtiter wells which contained factor VIII heavy chain that was immobilized using the anti-heavy chain monoclonal antibody ESH5 (black bars). To correct for background binding, phage were also incubated on microtiter wells that did not contain factor VIII heavy chain (white bars). On the y-axis the OD (450-540 nm) is depicted.

Figure 11A shows the deduced amino acid sequence of recombinant antibody fragments specific for the factor VIII heavy chain. The amino acid sequence of germ line variable heavy chain gene segments DP10 and DP47 is given. Deviations in amino acid sequence from these germline gene segments are indicated for two clones that encode recombinant antibodies that bind to the factor VIII heavy chain. Also the amino acid of the CDR3 and FR4 of the factor VIII heavy chain specific recombinant antibodies encoded by the two clones is given.

Figures 11B and C give the nucleotide and amino acid sequence of the variable heavy chain domain of two clones that encode recombinant antibodies that bind specifically to the factor VIII heavy chain.

DETAILED DESCRIPTION OF THE INVENTION

A number of investigators have addressed the epitope-specificity and mode of action of factor VIII inhibitory antibodies. Molecular cloning of the factor VIII cDNA revealed that factor VIII consists of a series of repeated domains which appear in the order A1-A2-B-A3-C1-C2. In plasma, factor VIII circulates as a heterodimer which consists of a heavy chain of variable length (90-220 kDa) and a light chain of 80 kDa. The factor VIII light chain consists of the domains A3-C1-C2 while the factor VIII heavy chain comprises the domains A1-A2-B. Heterogeneity of the factor VIII heavy chain is caused by limited proteolysis within the B-domain which contains several sites that are sensitive towards proteolytic cleavage. In plasma, factor VIII circulates in complex with von Willebrand factor, a large multimeric protein involved in the initial steps of platelet adhesion to a damaged vessel wall. Binding to von Willebrand factor protects factor VIII from proteolytic degradation. The physiological importance of this interaction is underscored by the low levels of factor VIII in plasma of patients that lack von Willebrand factor. Factor VIII is a precursor molecule which upon activation functions as a cofactor for factor IXa in the phospholipid and Ca^{2+} -dependent conversion of factor X to factor Xa. Activation of factor VIII involves proteolytic cleavages in both the heavy and light chain of factor VIII. Thrombin is considered to be the physiological activator of factor VIII and cleaves at Arg³⁷², Arg⁷⁴⁰ and Arg¹⁶⁸⁹ of factor VIII. Thrombin activated factor VIII thus consists of a hetero-trimer of the separate A1 and A2-domains and the cleaved factor VIII light chain (A3-C1-C2). Cleavage at Arg¹⁶⁸⁹ of the factor VIII light chain results in removal of amino-acid sequence Glu¹⁶⁴⁹-Arg¹⁶⁸⁹ which is essential for binding of factor VIII to von Willebrand factor. So far, three major binding sites for factor VIII inhibitors have been characterized (Scandella et al. 1994, Blood 86: 1811-1819; Healey et al. 1995, J. Biol. Chem. 270: 14505-14509; Fijnvandraat et al. 1998, Blood 91: 2347-2352).

Amino acid residues Val²²⁴⁸-Ser²³¹² in the C2-domain constitute a binding site for factor VIII inhibitors. The large size of this epitope suggests that a number of antibodies which bind to different amino acid regions in this area occur in plasma of patients with inhibitors of C2-specificity. The mechanisms of action of anti-C2 antibodies has been explored in considerable detail. Most of these antibodies interfere with binding of factor VIII to phospholipids. Furthermore, some of the antibodies with C2-specificity also inhibit the interaction of factor VIII with its carrier von Willebrand factor. A new

mechanism for inhibition of factor VIII by a human alloantibody has been described recently (Saenko et al. 1996, J. Biol. Chem. 271: 27424-27431). A human alloantibody that binds only to the amino-terminal portion (Val²²⁴⁸-Gln²²⁸⁵) of the C2-epitope has been shown to inhibit the thrombin induced release of factor VIII from von Willebrand factor.

Amino acid residues Arg⁴⁸⁴-Ile⁵⁰⁸ in the A2-domain of factor VIII constitute a major epitope for factor VIII inhibitors. Studies on the mechanism of inhibition of anti-A2 antibodies have shown that anti-A2 antibodies interfere with conversion of factor X to Xa by the lipid bound factor VIIIa-factor IXa-complex (Lollar et al., 1995). The anti-A2-antibodies do not interfere with binding of factor X to the factor VIIIa-factor IXa complex but simply limit the conversion of factor X.

A third major epitope of factor VIII inhibitors has been found in the A3-domain of factor VIII. Binding of inhibitory antibodies was dependent on the presence of amino acids Gln¹⁷⁷⁸-Met¹⁸²³. Previous studies have shown that this site constitute a binding site for factor IXa and indeed antibodies binding to this site interfered with complex assembly of factor VIIIa and factor IXa (Fijnvandraat et al. 1998, Blood 91: 2347-2352). In a number of patients with an inhibitor, inhibitory antibodies directed against other epitopes have been observed. An early study has shown that inhibitory antibodies may recognize amino acid region Met³³⁶-Arg³⁷² of factor VIII (Ware et al. 1988, Proc. Natl. Acad. Sci USA 85: 3165-3169). The mechanism of inhibition has not yet been explored but recently a binding site for factor X has been proposed in this part of the factor VIII molecule (Lapan, K.A. and Fay, P.J. 1997, J. Biol. Chem. 272: 2082-2088).

The restricted epitope specificity of factor VIII inhibitors suggests that a limited number of dominant B-cell epitopes is involved in the immune response to factor VIII. Apparently, human anti-factor VIII antibodies synthesized by B-cell clones from a variety of patients are surprisingly similar with respect to epitope specificity. This suggests that the primary amino acid and nucleotide sequence of antibodies with factor VIII specificity is similar at the molecular level. Based on this it is desirable to define the presence and epitope specificity of anti-factor VIII antibodies by simply addressing the presence of nucleotide sequences that correspond to antibodies with factor VIII inhibiting capacity. Sofar, the primary sequences of anti-factor VIII antibodies have been poorly defined. Davies and co-workers have suggested an association between factor

VIII inhibitors and use of VH gene segment DP73 (Davies et al. 1997, *Thromb. Haemostas.* supplement: 2352A). The nucleotide and primary amino acid sequence of these antibodies has not been disclosed and details with respect to the epitope specificity of these antibodies are lacking. Clearly, there is a need to
5 define the primary amino acid and nucleotide sequence of factor VIII antibodies in more detail. Such sequence information can be used to design diagnostic tests which can be used to monitor the occurrence of B-cell clones that produce factor VIII inhibitors in patients with haemophilia A. These diagnostic tests can be extremely sensitive and give information on the epitope specificity of factor VIII
10 inhibitors.

Studies directed at defining the epitope specificity and mode of action of these antibodies are limited by the heterogeneity of these antibodies in the plasma of these patients. Clearly, more stringent diagnostic criteria would be required to define the properties of factor VIII inhibitors in more detail.

15 A sudden increase in the frequency of inhibitor development in a group of previously treated patients has been associated with a particular pasteurized factor VIII concentrate manufactured in the Netherlands (Roosendaal et al. 1993, *Blood* 81: 2180-2186). These factor VIII inhibitors are directed against the factor VIII light chain and epitope mapping revealed that the majority of inhibitors
20 reacted with epitopes in the A3-C1 and the C2-domain of factor VIII (Sawamoto et al. 1998, *Thromb. Haemostas.* 79: 62-68). Recently, a second pasteurized factor VIII concentrate has been implicated in the development of inhibitors in a group of previously treated patients. Also in this case the inhibitory antibodies were predominantly of factor VIII light chain specificity (Peerlinck et al. 1997,
25 *Thromb. Haemostas.* 77: 80-86). It has been suggested that inhibitor development in these patients is due to small alterations in the factor VIII molecule which have been induced by the manufacturing process. This may indicate that the antibodies that developed in these patients have different properties compared to the factor VIII inhibitory antibodies that develop in other patients. Clearly,
30 knowledge of nucleotide and amino acid sequence of factor VIII specific antibodies could provide additional information on the etiology of factor VIII inhibitor which is desirable for the characterization of the antibody response in patients who have received these factor VIII concentrates.

Until now, the primary nucleotide and amino acid sequence of anti-factor
35 VIII antibodies has not been disclosed. This invention describes the nucleotide sequences that encode human antibodies with factor VIII-specificity. Based on

the primary sequence of these antibodies, oligonucleotide primers are designed that allow for detection of B-cells that produce antibodies with affinity for factor VIII. Detection of factor VIII specific B-cells may be accomplished using both mRNA, cDNA or DNA which are derived from lymphocytes of patients.

- 5 Genomic DNA, RNA and cDNA are prepared from lymphocytes by methods that are generally known in the art. Some methods for the detection of factor VIII specific B-cell clones are listed below. Other methods for the detection of nucleotide sequences of factor VIII specific antibodies, disclosed in this invention, are considered to fall within the scope of this invention. Selective
- 10 amplification of heavy chain variable sequences (VH-genes) can be used to detect nucleotide sequences that encode antibodies that are part of the human antibody repertoire that can bind specifically to factor VIII. The variable part of the human heavy chain is assembled from the variable heavy chain regions (VH), the diversity regions (D) and the joining regions (J). Fusion of these three
- 15 different gene segments is not a precise event and this so-called "junctional diversity", together with the process of nucleotide addition and deletion, results in the generation of the hypervariable complementary determining region 3 (CDR3). The human light chain is assembled in a similar manner but lacks diversity region D. Additional sequence diversity of both heavy and light chain
- 20 sequences is generated by somatic hypermutation and together with the mechanisms outlined above this ultimately results in the generation of high affinity antibodies. Knowledge on the nucleotide sequences that encode factor VIII-specific antibody allows for the detection of this specific antibody in the repertoire of patients who are at risk of developing factor VIII-specific antibodies
- 25 (such as haemophilia A patients who are treated with factor VIII or patients with acquired haemophilia). Amplification may be performed with a combination of oligonucleotide primers directed against constant regions or variable regions of heavy and light chain of factor VIII-specific antibodies. Detection of factor VIII specific antibodies may be performed using one oligonucleotide primer derived
- 30 from the variable parts of the nucleotide sequences encoding factor VIII antibodies and one oligonucleotide primer that is derived from the constant regions of factor VIII specific antibodies. Detection may also be performed using two oligonucleotide primers specific for variable parts of the nucleotide sequence that encodes an antibody that binds to factor VIII. The methods described herein
- 35 also include the amplification of immunoglobulin genes using oligonucleotide primers that are directed against the constant regions of the immunoglobulin

genes. Subsequent detection of nucleotide sequences of factor VIII specific antibodies can be performed using selective hybridization with (radiolabelled) oligonucleotide primers that are directed against the variable parts of the nucleotide sequence encoding factor VIII specific antibodies. From the above it follows that oligonucleotide primers are preferentially but not exclusively directed towards the constant and variable regions of factor VIII specific antibodies. In example 5, methods are disclosed that can be used to detect the presence of factor VIII specific antibodies in a mixture of nucleotide sequences. Combination of oligonucleotide primers derived from the nucleotide sequence of factor VIII specific antibodies can be used to directly assess the presence of factor VIII specific antibodies in the antibody-repertoire of patients. Alternatively, analysis by methods that include but are not limited to sequencing analysis, re-amplification of obtained fragments with more specific oligonucleotide primers, digestion with restriction enzymes and selective hybridization may be utilized to address the presence of factor VIII antibodies. Quantification of the amount of nucleotide sequences encoding factor VIII antibodies may be obtained by various methods that are generally known in the art and include but are not limited to the following. The amount of radioactivity incorporated into a PCR-fragment that encodes part of a factor VIII specific antibody can be determined. Furthermore, radioactively labelled oligonucleotide probes can be used to estimate the amount of a nucleotide sequence encoding a factor VIII specific antibody in a mixture of DNA fragments that code for part of a patients antibody repertoire. Quantitative PCR-amplification can be performed using for example dye-modified oligonucleotide primers which allow for direct monitoring of the amount of PCR-product generated during amplification.

Other methods that selectively detect and quantify specific nucleotide sequences that encode factor VIII specific antibodies may be devised by an average expert in the art. These methods are considered to fall within the scope of the present invention.

Examples 1-10 provide details on the identification and detection of nucleotide sequences that encode factor VIII specific antibodies in haemophilia patients. These examples teach how to arrive at the nucleotide sequence of factor VIII inhibitors and provide information on how to use this information for the detection of factor VIII specific antibodies.

This invention discloses the nucleotide and primary amino acid sequences of factor VIII specific antibodies. Factor VIII inhibitors are commonly directed

against three major epitopes on factor VIII within the A2- A3 and C2-domain of factor VIII. In Example 4 the nucleotide and amino acid sequence of anti-C2 antibodies is disclosed. In examples 8 and 9, the nucleotide and amino acid sequence of anti-A2 and anti-A3-C1 antibodies is disclosed. This invention teaches how to arrive at the nucleotide and amino acid sequence of factor VIII specific antibodies and the methods disclosed in this invention can be used to derive the nucleotide and amino acid sequence of anti-factor VIII antibodies with specificity for other domains of factor VIII which are a target for factor VIII inhibitors. Anti-factor VIII antibodies encoded by the nucleotide sequences disclosed here, can be used for the development of therapeutic agents that are capable of limiting the biological activity of factor VIII inhibitors. These therapeutic agents preferentially contain, but are not limited to:

1. The antibody fragments (or agents based on the nucleotide or amino acid sequence) can be used for the generation of anti-idiotypic antibodies. Antibody fragments can be either administered together with factor VIII or administered alone. Also, peptides or related agents which are based on the primary amino acid sequence of the variable parts of factor VIII specific antibodies can be used to induce the formation of anti-idiotypic antibodies direct against factor VIII inhibitors. Anti-idiotypic antibodies can also be obtained by screening large (semi-synthetic) libraries that encode a wide variety of recombinant antibodies. The preparation of anti-idiotypic antibodies can also take place in animals that include but are not limited to mouse by injection of recombinant antibody disclosed in this invention. Anti-idiotypic antibodies can subsequently be obtained by methods that are known to those skilled in the art. An anti-idiotypic response to factor VIII specific antibodies may also be obtained by injection of DNA encoding part of the nucleotide sequences of factor VIII specific antibodies which can be obtained by the methods outlined in this invention. Immunization by injection of DNA is considered to be only modestly immunogenic and other agents are needed to obtain a sufficiently high immune response. Co-injection of plasmid DNA encoding IL-2, GM-CSF and tetanus toxoid has been used to enhance the immune response to injected DNA (Spellerberg et al. 1997. J. Immunol. 15:1885-1892). Similar methods can be applied to enhance the immune response towards DNA fragments encoding factor VIII specific antibodies.

2. The recombinant antibody fragments described in this invention can be used as a therapeutic for treatment of patients with an inhibitor. Examples 7 and

10 disclose that recombinant antibody fragments (termed scFv's) which bind specifically to the C2-domain, interfere with binding of inhibitory antibodies to factor VIII. These scFv's can be used for treatment of patients with inhibitory antibodies directed against the C2-domain. This invention discloses how to arrive
5 at recombinant antibody fragments that bind specifically to the A2, A3-C1 and C2-domain of factor VIII. Using the methods outlined in this invention additional antibody fragments directed against these and other regions on factor VIII may be obtained. These regions include but are not limited to Arg⁴⁸⁴-Ile⁵⁰⁸ in the A2-domain, Gln¹⁷⁷⁸-Met¹⁸²³ in the A3-domain and Val²²⁴⁸-Ser²³¹² in the C2-domain.
10 Recombinant antibodies directed against multiple epitopes preferentially will be part of a pharmaceutical preparation since most patients have inhibitory antibodies directed against multiple epitopes on factor VIII. The inhibitor neutralizing activity of the recombinant antibody fragments described in this invention may be modified by the introduction of point mutations in the constant
15 and variable parts of these fragments. Furthermore, the recombinant antibody fragments described in this invention may be cloned into vectors which allow for expression of these fragments for example as Fab-fragment. Other vectors for expression of antibodies and antibody fragments are available to an average expert in the field (see for example: "Antibody Engineering; A Practical
20 Approach" edited by Mc Cafferty et al. 1996. Oxford University Press). Methods to increase the affinity of recombinant antibodies or antibody fragments are readily available and can be used to modulate the biological activities of the recombinant antibody fragments described in this invention. Peptides and peptide-related agents which have been designed based on the amino acid
25 sequence of the recombinant antibodies described in this invention (for example peptides derived from the amino acid sequence of CDR3) can be used to interfere with the activity of factor VIII inhibitors.

In summary, this invention provides an improved method to define factor VIII inhibitory antibodies. The methods described result in definition of the
30 nucleotide sequence of factor VIII inhibitors and teaches how to utilize this information for the development of improved methods for the detection of factor VIII inhibitors. This invention also discloses pharmaceutical preparations, derived from the nucleotide and/or primary amino acid sequence of factor VIII specific antibodies, that can be used in the treatment of haemophilia A patients
35 with inhibitory antibodies directed against factor VIII. The dose of the therapeutic agents to be administered to a patient, obviously depends on the

affinity of the therapeutic agent for factor VIII. The affinity of the recombinant antibody fragments described in this invention can vary considerably as is disclosed in Examples 6 and 7. The dose to be administered also depends on bodyweight of the patient, the titre of the factor VIII inhibitor and the biological activities of the different components of the therapeutic agent used. The dose to be administered can be estimated according to methods that are disclosed in Examples 7 and 10. These methods may be complemented with dose finding studies which involve administration of the therapeutic agents in animal models and healthy individuals. In general, the dose administered will vary between 10 μ g - 5 g/kg and more preferably between 100 μ g - 1g/kg of bodyweight per day.

The therapeutic agent may be administered in combination with factor VIII (or a substitute of factor VIII). The dose of factor VIII administered together with the therapeutic agent may vary between 0.1 and 2000 Units per kg and more preferably between 1 and 200 Units per kg of body weight per day.

The words "substantially isolated form" as used herein are intended to refer to a form of the material which is different from any naturally occurring form of said material, i.e. different from the material in its natural environment. In particular, the words are intended to define relative freedom or absence of substances that naturally accompany the material.

EXAMPLE 1: Characterization of anti-factor VIII antibodies in patient's plasma

Anti-factor VIII antibodies present in the plasma of a patient with acquired haemophilia were characterized by immunoprecipitation and neutralization experiments. The construction of recombinant factor VIII fragments corresponding to the A2, A3-C1-C2 and C2-domain of factor VIII has been described previously (Fijnvandraat et al. 1997. Blood 89: 4371-4377; Fijnvandraat et al. 1998. Blood 91: 2347-2352). These recombinant factor VIII fragments were metabolically labelled with [35 S]-methionine and subsequently used for the detection of anti-factor VIII antibodies by immunoprecipitation using methods that have been described previously (Fijnvandraat et al. 1998. Blood 91: 2347-2352). Reactivity with both metabolically labelled A2, A3-C1-C2 and C2 domain was observed (data not shown). This indicates that at least two classes of antibodies directed against factor VIII were present in the plasma of the patient. To determine the contribution of the different antibodies in the patient's plasma to the titre of the inhibitor as measured in the Bethesda assay we

performed neutralization experiments. Increasing concentrations of recombinant factor VIII fragments were mixed with samples that contained factor VIII antibodies diluted until a final inhibitory capacity of 2 BU/ml. Addition of both recombinant factor VIII light chain (A3-C1-C2) and C2-domain resulted in a decrease in the inhibitory activity of 50 and 20%, respectively. Addition of the factor VIII heavy chain (domains A1-A2-B) resulted in 45% neutralization of the inhibitor in the plasma of the patient. Based on these data we conclude that inhibitory antibodies directed against the heavy chain contribute for 45% to the inhibitory capacity of the patient's anti-factor VIII antibodies whereas anti-A3-C1-C2 antibodies account for the other half of the inhibitory capacity. Our results provide evidence for the occurrence of at least three classes of inhibitory antibodies in the patient's plasma. Next, we determined the subclass of the anti-factor VIII antibodies using methods that have been outlined previously (Fijnvandraat et al. 1997. Blood 89: 4371-4377). The antibodies with A2-specificity consisted predominantly of subclass IgG4; in addition small amounts of subclass IgG2 were observed. The antibodies directed against the factor VIII light chain consisted exclusively of subclass IgG4. The methods outlined above provide a starting point for further characterization of human antibodies with specificity for factor VIII. Similar analyses can be performed on samples derived of other patients which are analyzed for the presence of factor VIII inhibitors.

EXAMPLE 2: Construction of an IgG4 specific library

Peripheral blood lymphocytes were isolated from a blood sample of a patient with acquired haemophilia. The titre of the inhibitor was 1250 BU/ml. RNA was isolated from the lymphocytes using RNazol (WAK Chemie, Germany) according to the instructions of the manufacturer. RNA was transcribed into cDNA employing random hexamer primers (Gibco, Breda, The Netherlands). Since, most of the anti-factor VIII antibodies described in Example 1 were of subclass IgG4, DNA fragments corresponding to the heavy chain of immunoglobulins of subclass IgG4 were amplified using the following set of oligonucleotide primers:

```

conIgG1-4   5' CTTGTCCACCTTGGTGTGCTGGG 3'
huIgG4      5' ACGTTGCAGGTGTAGGTCTTC   3'
35 huVH1aback 5' CAGGTGCAGCTGGTGCACTCTGG 3'
huVH2aback  5' CAGGTCAACTTAAGGGAGTCTGG 3'

```

	huVH3aback	5'	GAGGTGCAGCTGGTGGAGTCTGG	3'
	huVH4aback	5'	GAGGTGCAGCTGTTGCAGTCGGG	3'
	huVH5aback	5'	GAGGTACAGCTGCAGCAGTCTGC	3'
	huVH6aback	5'	GAGGTACAGCTGCAGCAGTCTGC	3'
5	huJH1-2forSal	5'	GAGTCATTCTCGTGT <u>TCGAC</u> ACCGGTGACCAGGGTGCC	3'
	huJH3forSal	5'	GAGTCATTCTCGTGT <u>TCGAC</u> ACCGGTGACCATTGTCCC	3'
	huJH4-5forSal	5'	GAGTCATTCTCGTGT <u>TCGAC</u> ACCGGTGACCAGGGTTCC	3'
	huJH6forSal	5'	GAGTCATTCTCGTGT <u>TCGAC</u> ACCGGTGACCCTGGTCCC	3'
	huVH1backNco	5'	AATCCATGGCCAGGTGCAGCTGGTGCA	3'
10	huVH2backNco	5'	AATCCATGGCCAGGTCAACTTAAGGGA	3'
	huVH3backNco	5'	AATCCATGGCCAGGTGCAGCTGGTGGA	3'
	huVH4backNco	5'	AATCCATGGCCAGGTGCAGCTGTTGCA	3'
	huVH5backNco	5'	AATCCATGGCCAGGTACAGCTGCAGCA	3'
	huVH6backNco	5'	AATCCATGGCCAGGTACAGCTGCAGCA	3'

15

Oligonucleotide primers huVHa(1-6)back and huJH(1-6)forSal have been described previously (Marks et al. 1991, J. Mol. Biol. 222: 581-597). Oligonucleotide primers huVH(1-6)backNco have been adapted from oligonucleotide primers described in the same paper. The first series of amplification involved primers huVH(1-6)back in conjunction with primer conIgG1-4. Six different DNA fragments of about 700 bp, each corresponding to an individual VH-gene family were obtained. The six different fragments were isolated and re-amplified with primers huVH(1-6)back and primer huIgG4. Six products of approximately 660 bp were obtained. The 6 different 660 bp fragments which represented the IgG4 repertoire of the patient were re-amplified with primers huVH(1-6)backNco and huJH(1-6)forSal in order to prepare these fragments for cloning. The resulting 24 fragments were pooled according to VH-gene family and the six different fragments were digested with NcoI and SalI. The digested fragments were purified and dissolved in TE (10 mM Tris-HCl pH=8.0; 0.1 mM EDTA). The vector pHEN-1-VLrep has been described previously (Griffin, H.M. and Ouwehand, W.H. 1995. Blood 86, 4430-4436; Schier et al. 1996. J. Mol. Biol. 255: 28-43) and contains a light chain repertoire derived of two non-immunized donors. Insertion of a heavy chain repertoire in this vector has been shown to result in the production of antibody fragments that consist of the variable domains of both heavy and light chain. These antibody fragments have been termed single chain Fv (scFv) fragments (Hoogenboom,

35

H.R. et al. 1991. Nucleic Acid Res. 19: 4133-4137). The vector pHEN-1-VLrep (kindly provided by Dr. W.H. Ouwehand, Department of Transfusion Medicine, University of Cambridge, UK) was digested with XhoI and NcoI and the six fragments corresponding to the IgG4-specific heavy chain repertoire of the patient with acquired haemophilia were inserted. The ligation mixtures were transformed to *E. coli* TG1 and a library of 1.500.000-2.500.000 independent clones was obtained. Colonies were scraped and resuspended in 2TY supplemented with 15% glycerol, 100 µg/ml ampicillin and 1% glucose. Similar to the methods outlined above libraries that represent the immunoglobulin repertoire of other patients may be assembled.

EXAMPLE 3: Selection of factor VIII specific antibodies

Selection of clones that encoded antibody fragments (scFvs) with factor VIII specificity was performed as outlined below. Glycerol stocks were plated onto 2TY plates that contained ampicillin (100 µg/ml) and 1% glucose. Colonies were grown overnight and scraped the next day and dissolved in 2TY supplemented with 100 µg/ml ampicillin and 1% glucose. These cells were diluted in 2TY supplemented with ampicillin (100 µg/ml) and 1% glucose till a final optical density (OD) of 0.3 (measured at 600 nm). Cells were grown at 37°C till an OD of 0.5. Subsequently, 1 ml of culture was diluted 10 times in 2TY with ampicillin (100 µg/ml) and 1% glucose. Next, 20 µl of helper phage was added (VCSM13; 1×10^{11} pfu/ml) and the mixture was incubated for 45 minutes at 37°C without shaking. Then, cells were incubated at 37°C with shaking at 150 rpm for another 45 minutes. The cells were spun down at low speed and resuspended in 100 ml of 2TY supplemented with ampicillin (100 µg/ml), 0.1% glucose and 25 µg/ml kanamycin. The cells were incubated overnight at 30°C. The next day cells were spun down at 10000 rpm for 30 minutes. The supernatant was harvested and recombinant phage were allowed to precipitate for 2 hours at 4°C after the addition of 1/5 volume of 20% PEG6000/2.5 M NaCl. The phages were spun down (30 minutes 10000 rpm) and resuspended in 5 ml of TBS (50 mM Tris-HCl pH 7.4, 150 mM NaCl). This preparation was spun down for 5 minutes at 14000 rpm and the supernatant was stored at 4°C.

Selection of factor VIII-binding phages was performed as outlined below. Microtiter wells were coated overnight at 4°C with 5 µg/ml of the murine factor VIII light chain specific monoclonal antibody CLB-CAG 12 which was diluted in 50 mM NaHCO₃ (pH 9.5). The wells were blocked for 1 hour at 37°C with TBS

containing 3% HSA. Phage solution (1×10^{12} pfu/ml) was diluted 1 to 1 in TBS supplemented with 6% HSA and 1% Tween-20 and incubated for 2 hours at room temperature with microtiter wells that contained immobilized CLB-CAg 12. The phage solution was removed and transferred to a second microtiter well (also coated with CLB-CAg 12) which had been preincubated with 1 μ g/ml of factor VIII light chain. Phages were incubated with factor VIII light chain for 2 hours at room temperature. Wells were washed 20 times with TBS/0.1% Tween-20 and 20 times with TBS and bound phage was eluted with 100 mM triethylamine (pH 12). Eluted phage (volume 1 ml) was neutralized by the addition of 500 μ l of 1 M Tris-HCl pH 7.4 and subsequently added to 5 ml of *E. coli* TG1 (OD 600 = 0.5). Cells were incubated for 1/2 hour at 37°C (no shaking) and 10 minutes at 37°C (shaken at 200 rpm). Cells were collected by centrifugation for 10 minutes at 4000 rpm for 7 minutes. Subsequently, infected TG1 cells were plated on 2TY agar plates supplemented with ampicillin (100 μ g/ml) and 1.0% glucose. Cells were grown overnight at 30°C. Cells were scraped in 2TY supplemented with ampicillin (100 μ g/ml), 1% glucose and 15% glycerol. Ampoules were stored at -70°C.

Alternatively, factor VIII light chain (5 μ g/ml in 50 mM NaHCO₃ (pH 9.6)) was immobilized on immunotubes (Nunc, Life Technologies, Breda, The Netherlands). First, 1 ml of phage solution diluted in TBS supplemented with 3% HSA was incubated for 2 hours at room temperature in non-coated Immunotubes. Subsequently, 1 ml of phage solution was removed and incubated for 2 hours at room temperature in immunotubes coated with factor VIII light chain. Immunotubes were washed 20 times with TBS/0.1% Tween-20 and 20 times with TBS. Bound phage was eluted with 100 mM triethylamine (pH 12) and processed as outlined above.

The second round of panning was initiated by inoculating 50 μ l of glycerol stock obtained after the first selection in 10 ml 2TY supplemented with ampicillin (100 μ g/ml) and 1% glucose till a final OD600 of 0.3. Cells were grown till an OD600 of 0.5, diluted 1 to 10 in 2TY supplemented with ampicillin (100 μ g/ml), kanamycin (25 μ g/ml) and 1% glucose and subsequently infected with a 20 fold excess of VCSM13. Cells were grown overnight at 30°C. Supernatant containing the phage was harvested as described above and again screened for binding to immobilized factor VIII light chain. After four rounds of panning, 30 clones selected by ELISA and 30 clones that were obtained after panning with factor VIII light chain immobilized to immunotubes were grown

and analyzed for binding to the factor VIII light chain. Colonies were picked and grown overnight in 2 ml 2TY supplemented with 100 µg/ml ampicillin and 1% glucose. The next day the cultures were diluted 200 times and grown till an OD₆₀₀ of approximately 0.5. Cells were subsequently infected with VCSM13
5 for 45 minutes at 37°C (no shaking) and 45 minutes at 37°C (shaken at 200 rpm). Infected cultures were diluted 1 to 10 in 2TY supplemented with 100 µg/ml ampicillin, 0.1% glucose and 25 µg/ml kanamycin and cells were grown overnight at 30°C. Supernatant containing phage was collected after centrifugation and tested for binding to the factor VIII light chain as described below.
10 The non-inhibiting murine monoclonal antibody CLB-CAG 12 directed against an epitope in the A3-C1 domain of factor VIII was immobilized on microtiter wells at a concentration of 5 µg/ml in 50 mM NaHCO₃ (pH 9.5). Wells were blocked for 1 hour in TBS supplemented with 3% HSA. Wells were incubated with factor VIII light chain at a concentration of 1 µg/ml in 50 mM Tris HCl (pH
15 7.4), 1 M NaCl, 2% HSA for 2 hours at 37°C. Fifty µl of phage solution and an equal volume of TBS supplemented with 1% Tween-20 and 6% HSA were added to wells containing factor VIII light chain. To monitor specific binding of the phages, wells that did not contain factor VIII light chain were incubated simultaneously with the same phage solution. Phage was incubated at room temperature for 2 hours and were shaken at regular intervals. Subsequently, wells
20 were extensively washed 5 times with TBS supplemented with 0.1% Tween-20 and washed 5 times with TBS. The presence of bound phage was monitored by incubating with a peroxidase labelled polyclonal antibody directed against M13 (Pharmacia-LKB, Woerden, The Netherlands) in a dilution of 1 to 4000 in TBS
25 supplemented with 1% HSA and 0.1% Tween-20. Plates were washed 5 times with TBS supplemented with 0.1% Tween-20 and 5 times with TBS. Binding of peroxidase-labelled anti-M13 antibody was quantified by incubation with 3-3'-5-5' tetramethylbenzidine (TMB). Substrate conversion was arrested by the addition of 100 µl of 2N H₂SO₄. Part of the results of this analysis are given in
30 Figure 1. An example of 12 clones that show specific binding to the factor VIII light chain is given. Clearly, phage encoded by clone 1 to 12 display binding to the factor VIII light chain (black bars). Some background binding is visible which is not dependent on the presence of the factor VIII light chain (grey bars). The bars labelled with c represent two clones that express antibody fragments
35 that do not bind specifically to the factor VIII light chain. These clones have been derived from the initial library and have not been selected on the factor VIII light

chain. To ensure that during subsequent rounds of panning an increase in the amount of factor VIII specific recombinant antibody fragments was obtained, we screened 12 clones obtained after the first round of panning for binding to the factor VIII light chain (Figure 2). Only 3 out of 12 clones bind specifically to the factor VIII light chain. In 9 out of 12 clones binding of phage is not dependent on the presence of the factor VIII light chain. These results clearly indicate that during panning the amount of phages that express factor VIII-specific antibodies can be selectively enriched. In summary, we have outlined a specific protocol for the selection of factor VIII specific antibodies that correspond to the spectrum of anti-factor VIII antibodies present in the patient with acquired haemophilia. In the first two examples our analysis is limited to material derived of one single patient and only antibodies directed against the factor VIII light chain have been analyzed. Using the methods outlined in these two examples the repertoire of anti-factor VIII antibodies of other patients with an inhibitor can easily be obtained. Furthermore, anti-factor VIII antibodies directed against epitopes located outside the factor VIII light chain may be obtained by adapting the screening methods used in Example 1 and 2. Antibodies directed against the heavy chain can be selected by immobilizing factor VIII heavy chain employing monoclonal antibody CLB-Cag 9. In these two examples we have focused on the IgG4-repertoire of the patient. Similarly, other subclasses may be investigated using the appropriate primers. For example, subclass IgG1-4 can be detected by simply using primer conIgG1-4 described in Example 1. Similarly, other primers specific for IgA, IgM, IgE and IgD may be utilized to assemble antibody-repertoires that include factor VIII-specific antibodies.

EXAMPLE 4: Sequence characteristics of recombinant antibodies with factor VIII light chain specificity

In the previous examples methods to obtain recombinant antibodies with factor VIII specificity has been outlined. To obtain information on the properties of these antibodies we selected 30 clones that have been selected by immobilized factor VIII in immunotubes. Also 30 clones which were selected employing factor VIII light chain with monoclonal antibody CLB-Cag 12 were analyzed. Clones were grown as described in Example 2 and plasmid DNA was isolated. The nucleotide sequence of the variable part of the heavy chain (VH domain) of 55 clones was determined using fluorescently labelled M13 reverse primer on an ABI-Prism 377 DNA sequencer. The sequences obtained were aligned with

heavy chain sequences in the database "V BASE" of the MRC Centre of Protein Engineering (Cambridge, UK). The 55 clones analyzed were encoded by two different VH-gene segments DP-10 and DP-14 (Cook and Tomlinson, Immunology Today 16: 237-242). The 41 clones that were encoded by the germline sequence DP14 consisted of three groups of recombinant antibodies that differed mainly in the nucleotide sequences of the constant regions of the VH gene. Thirty-three clones which were represented by clone IT2, 5 clones were represented by EL25 and 3 clones were represented by clone EL5 (Table I). Two clones that were encoded by DP10 (EL14) and DP14 (IT2) were selected for further analysis. The nucleotide and primary amino acid sequence of these clones is listed in Figure 3 and 4. The characteristics of the two sequences are given in Table I. Part of clone EL14 is most likely derived of the D-segment D6-13 and J-segment JH-3b. Somatic hypermutation has occurred during the immune response as evidenced by the large number of nucleotide changes compared to the germline sequences of the VH segments. The variable heavy chain part of clone IT2 contains 20 nucleotide substitutions when compared to the germ line segment DP-14. These 20 nucleotide substitutions result in a total of 13 amino acid changes (Table I). The variable heavy chain part of clone EL14 contains 18 nucleotide substitutions when compared to the germ line segment DP-10. These 18 nucleotide substitutions result in 12 amino acid changes (Table I). Clone IT2 has in part been derived from gene segments D3-3 and JH6b. Remarkably, a stretch of G-residues is observed between the germ line sequences DP14 and D3-3 for clone IT2 that encodes for a flexible arm of glycine residues. Inspection of the amino acid sequence of clone EL14 and IT2 reveals several interesting features. Both CDR3 regions contain several glycine residues at their amino-terminal part which is in both cases followed by a tyrosine and a glutamic acid (GG-YE). Furthermore, a proline, alanine and an aspartic acid appear to be conserved in the carboxyl-terminal part of the CRD3 (P---A-D). A common motif can be derived from the amino acid sequences of the CDR3 regions of clone EL14 and IT2 which is given in Figure 4B. These features may determine the specificity of these antibodies for the factor VIII light chain. In this example the nucleotide and primary amino acid sequence of two recombinant factor VIII antibodies has been disclosed. With methods similar to the ones described in this example recombinant antibodies that are directed against other regions on the factor VIII molecule may be analyzed. Common features of these antibodies can be identified as outlined in this example and therapeutic and diagnostic agents

derived of these common features can be used for diagnosis and treatment of patients with factor VIII inhibitors.

EXAMPLE 5: Detection of nucleotide sequences of factor VIII specific antibodies in patient samples

The nucleotide and amino acid sequences outlines in the previous example can be used to specifically detect factor VIII antibodies with C2-specificity in heterogeneous mixtures of antibodies. This can be accomplished by developing reagents, for example, antibodies that specifically recognize the anti-factor VIII antibodies described in this invention. Detection of factor VIII-specific antibodies can also be performed by analysis of the presence of specific nucleotide sequences that encode factor VIII specific antibodies. Methods to obtain nucleotide sequences that encode factor VIII specific antibodies are disclosed in this invention. In this example the detection of nucleotide sequences encoding one of the factor VIII specific antibodies described in the previous example (EL14) is disclosed. Lymphocytes of the patient with acquired haemophilia described in the first example were obtained. RNA was isolated and cDNA was prepared. Subsequently, DNA fragments were amplified with oligonucleotide primers huVH(1-6)aback and conIgG1-4 (see Example 2). The six different 700 bp fragments obtained were isolated and used for a second PCR with oligonucleotide primer huVH(1-6)aback and huIgG4. This resulted in a fragment of 660 bp which was cloned into the vector pGEM-T (Promega, Madison, WI, USA). The presence of nucleotide sequences that corresponded to that of clone EL14 was addressed by nucleotide sequencing. One out of sixty clones analyzed did contain nucleotide sequences that were identical to that obtained for clone EL14. This analysis shows that, using the nucleotide sequences disclosed in this invention as a starting point, it is possible to monitor the presence of factor VIII specific antibodies in patient samples. In this example oligonucleotide primers are used which have also been employed for the construction of the IgG4 specific library. Other combinations of oligonucleotide primers that are based on the nucleotide sequences of clone EL14 and IT2 may be designed which may include but are not limited to oligonucleotide primers that are based upon the CDR3 region of these antibodies. In this example detection of factor VIII specific antibodies is performed using analysis of nucleotide sequences. Alternatively, detection of factor VIII specific sequences may also be performed employing selective hybridization using probes that are

based on the nucleotide sequence of the factor VIII specific antibodies disclosed in this invention. Other means of detection of specific nucleotide sequences that are known to an average expert in the art also fall within the scope of this invention. The methods disclosed in this invention allow for the isolation of factor VIII antibodies and determination of their nucleotide and amino acid sequence. In this example we have outlined described methods that detect factor VIII specific antibodies present in the repertoire of a patient with a factor VIII inhibitor. In examples 8 and 9 the nucleotide sequence of antibody fragments that bind to the A2- and A3-C1 domain of factor VIII is given. Methods similar to the ones described in this example can be used to detect nucleotide sequences that encode factor VIII inhibitors with A2-, A3-C1- or with a different epitope-specificity.

EXAMPLE 6: Properties of factor VIII-specific antibodies scFv-IT2 and scFv-EL14

The biochemical properties of the factor VIII specific antibodies IT2 and EL14 were characterized as follows. First, the plasmids pHEN-1-VL-EL14 and pHEN-1-VL-IT2 were digested with NcoI and NotI and the recombinant antibody fragments were isolated and cloned into the vector pUC119-sfi/Not-His6 (kindly provided by Dr. W.H. Ouwehand, University of Cambridge, Division of Transfusion Medicine, Cambridge UK). Positive clones were identified and grown till OD600 of 0.8-1.0 in 2TY medium supplemented with 1% glucose and 100 µg/ml ampicillin. Subsequently, Isopropyl-β-D-thiogalactopyranoside (IPTG) till a final concentration of 1 mM was added and cells were grown for 3 hours at 30°C. Cells were harvested by centrifugation for 15 minutes at 4000 g at 4°C. The pellet was dissolved in 10 ml of 30 mM Tris-HCl (pH 8.0), 1 mM EDTA and 20% sucrose in order to release the content of the periplasma. The mixture was incubated at 4°C for 20 minutes and subsequently cells were collected by centrifugation (15 min 10800 g at 4°C). The supernatant which consists primarily of proteins present in periplasma was collected. The pellet was resuspended in 10 ml 5 mM MgSO₄ and incubated for 20 minutes at 4°C. Residual cell debris was collected by centrifugation for 15 minutes at 10800 g. The supernatant (designated osmotic shock fraction) was collected and added to the fraction containing periplasma-derived proteins. The pooled fractions were centrifuged for 20 minutes at 30000 g at 4°C and the supernatant was collected. The supernatant was filtered over a 0.22 µm filter. A

mixture of protease inhibitors was added (Complete™ Mini, Boehringer Mannheim, Germany) and the pooled fractions were dialysed overnight against a buffer containing 50 mM NaPi (pH 7.4), 20 mM imidazole and 500 mM NaCl. ScFvs were purified by nickel affinity resin Ni-NTA (QIAGEN, Germany) as follows: 1 ml of Ni-NTA matrix was equilibrated with 50 mM NaPi (pH 7.4), 250 mM imidazole, 500 mM NaCl and subsequently with 50 mM NaPi (pH 7.4), 20 mM imidazole, 500 mM NaCl. Dialysed supernatant containing factor VIII specific scFv's were then batch-wise incubated with Ni-NTA matrix for 3 hours at 4°C. The Ni-NTA was then transferred to a column and washed with 7 ml of 50 mM NaPi (pH 7.4), 20 mM imidazole, 500 mM NaCl and 7 ml of 50 mM NaPi (pH 7.4), 35 mM imidazole, 500 mM NaCl. ScFv's were eluted with 50 mM NaPi (pH 7.4), 250 mM imidazole, 500 mM NaCl and stored at 4°C. The purity of the different scFv preparations was addressed by SDS-PAGE followed by staining with Coomassie Brilliant Blue. All purified ScFv's appeared for at least 90% homogenous and migrated with an apparent molecular weight of 30 kDa. In the preparations obtained a small amount of a protein with a lower molecular weight was observed. The identity of this band was investigated by immunoblotting with monoclonal antibody 9E10. The epitope of this antibody is present at the carboxyl-terminus of the scFv's. Both the protein migrating at a molecular weight of 30 kDa and 15 kDa reacted with monoclonal antibody 9E10 on Western blot. This indicates that the 15 kDa fragment most likely corresponds to the light chain of the scFv's. Purified scFv's corresponding to clone IT2 and EL14 were purified as outlined above. A scFv derived of clone O4, a clone present in the patient library that did not bind to the factor VIII light chain was included as a negative control. The binding of scFv-IT2 and scFv-EL14 to the factor VIII light chain was addressed employing the following ELISA. Monoclonal antibody 9E10 (5 µg/ml) dissolved in 50 mM NaHCO₃ pH 9.5 was immobilized on microtiter wells overnight at 4°C. Subsequently, purified scFv's diluted in 50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1% HSA and 0.2% Tween-20 were added and incubated for 2 hours at room temperature. The microtiter plates were washed 5 times with TBS, 0.1% Tween-20. Next, purified factor VIII light chain was added (5 µg/ml) together with peroxidase labelled monoclonal antibody CLB-Cag A (0.5 µg/ml). The mixture (diluted in TBS, 0.1% Tween-20) was incubated for 2 hours at room temperature. The microtiter wells were washed 5 times with Tris-buffered saline (TBS) supplemented with 0.1% Tween-20 and 5 times with TBS. The amount of bound peroxidase labelled monoclonal

antibody CLB-Cag A was quantified by the substrate TMB. The results of this analysis are given in Figure 5. Both scFv-EL14 and scFv-IT2 react with specifically with the factor VIII light chain while scFv-04 did not react with the factor VIII light chain. Next, we used the factor VIII inhibitory murine monoclonal antibody CLB-Cag 117 for the detection of bound factor VIII light chain. The ELISA was performed as outlined above. Instead of peroxidase labelled CLB-Cag A we used peroxidase labelled CLB-Cag 117 for the detection of immobilized factor VIII light chain. We did not observe binding of CLB-Cag 117 when factor VIII light chain is immobilized by scFv-EL14 and scFv-IT2 (Figure 5). These results show that the epitope of scFv-EL14 and scFv-IT2 overlaps with that of CLB-Cag 117. Previously, we have shown that the inhibitory antibody CLB-Cag 117 is directed against the C2-domain of factor VIII (Fijnvandraat et al. 1998. Blood 91: 2347-2352). Apparently, both scFv's bind to an epitope in the C2-domain of factor VIII which overlaps with that of the inhibitory murine monoclonal antibody CLB-Cag 117.

Next, different dilutions of scFv-EL14 and scFv-IT2 were tested for binding to immobilized factor VIII light chain as outlined above using CLB-Cag A as the detecting antibody (Figure 6). From this analysis it appeared that scFv-EL14 binds with a higher affinity to the factor VIII light chain than scFv-IT2. These results were complemented by immunoprecipitation experiments for scFv-EL14. Immunoprecipitation experiments employing a metabolically labelled fragment corresponding to the C2-domain was performed essentially as described previously (Fijnvandraat et al. 1998. Blood 91: 2347-2352). Monoclonal antibody 9E10 was covalently linked to CNBr-activated Sepharose 4B and this matrix was used to bind scFv-EL14. Specific binding of scFv-EL14 to metabolically labelled C2-domain was detected and this confirms the C2-specificity of this recombinant antibody fragment. In this example methods have been disclosed to characterize recombinant antibodies with specificity for the C2-domain. In examples 8 and 9, we describe the nucleotide and amino acid sequence of recombinant antibody fragments that bind specifically to the A2- and A3-C1 domain of factor VIII. The methods described in this example can easily be adapted by an average expert skilled in the art, which will allow for characterization of recombinant antibodies directed against the A2, A3-C1 or another epitope on factor VIII.

EXAMPLE 7: Factor VIII specific recombinant antibody fragments scFv-IT2 and scFv-EL14 neutralize the activity of factor VIII inhibitors

In the previous example, we have shown that scFv-EL14 and scFv-IT2 bind to the factor VIII light chain and compete for binding with the murine inhibitory monoclonal antibody CLB-CAg 117. These observations suggest that the epitope of both scFv-EL14 and scFv-IT2 overlaps with that of CLB-CAg 117. It is expected that similar to CLB-CAg 117, scFv-EL14 and scFv-IT2 inhibit the biological activity of factor VIII. Increasing amounts of purified scFv's were tested for inhibition in the Bethesda assay. Surprisingly, addition of up to 170 µg/ml scFv did not result in factor VIII inhibition as measured in the Bethesda assay. In contrast, CLB-CAg 117 readily inhibited factor VIII when measured in the same assay. Apparently, binding of scFv-EL14 and scFv-IT2 to factor VIII does not interfere with the biological activity of factor VIII. This finding prompted us to investigate the capacity of both scFv-EL14 and scFv-IT2 to overcome inhibition by CLB-CAg 117. Monoclonal antibody CLB-CAg 117 was diluted till a final inhibitory activity of 2 BU/ml. This value corresponds with a residual factor VIII activity of 25% in the Bethesda assay. Subsequently, increasing concentrations of scFv-EL14 and scFv-IT2 were added. Surprisingly, both scFv-EL14 and scFv-IT2 could overcome the factor VIII inhibitory activity of CLB-CAg 117 (Figure 7). ScFv-EL14 (panel A) proved to be more efficient than scFv-IT2 (panel B) in neutralizing the inhibitory activity of CLB-CAg 117. Both scFv-EL14 and scFv-IT2 were unable to neutralize the inhibitory activity of monoclonal antibody CLB-CAg A, directed against amino acid residues Glu¹⁸¹-Lys¹⁸¹⁸ on the factor VIII light chain (Lenting et al. 1996. J. Biol. Chem. 271: 1935-1940). These results for the first time show that antibody fragments with factor VIII specificity can be used to interfere with the activity of factor VIII inhibitors. Administration of these antibody fragments will be beneficial for the treatment of patients with inhibitory antibodies directed against factor VIII. In this example the biological activity of antibody fragments with C2-specificity is disclosed. In examples 8 and 9, the nucleotide and amino acid sequence of recombinant antibody fragments that bind to the A2 and A3-C1 domain of factor VIII is disclosed. The methods disclosed in this and the previous example can easily be adapted by an average expert skilled in the art to establish the capacity of recombinant antibody fragments directed against the A2 or A3-C1 domain to neutralize factor VIII inhibitors. Similar to outlined in this example recombinant antibody fragments that bind to other regions can be evaluated for their

neutralizing capacity of factor VIII inhibitors. Similarly to what has been described in this example for scFv-EL14 and scFv-IT2, antibody fragments binding to A2, A3-C1 and other domains on factor VIII can be used for treatment of patients with factor VIII inhibitors.

5

EXAMPLE 8: Isolation and characteristics of anti-factor VIII antibodies that specifically bind to the A3-C1 domain of factor VIII.

Previous studies have indicated that plasma of a substantial number of inhibitor patients contains anti-factor VIII antibodies that bind specifically to the A3-C1 domains of factor VIII (Fijnvandraat et al. 1998. Blood 91: 2347-2352; Zhong et al. 1998. Blood 92: 136-142). Here, we have employed phage display technology to isolate anti-factor VIII antibodies from the total immunoglobulin repertoire of a haemophilia A patient with an inhibitor. Previously, we have shown that in plasma of this patient anti-factor VIII antibodies directed against the factor VIII light chain are present. The majority of the anti-factor VIII antibodies in this patient is directed against the A3-C1 domain whereas a small portion of anti-factor VIII antibodies reacts with the C2-domain (Fijnvandraat et al. 1998. Blood 91: 2347-2352). The majority of anti-factor VIII antibodies were of subclass IgG4. An IgG4-specific library was constructed using peripheral blood lymphocytes of the patient as starting material. A library consisting of 1.9×10^6 independent clones was obtained using the methods outlined in Example 2. Selection of recombinant phage that bind specifically to factor VIII was performed essentially as outlined in Example 3.

Microtiter wells were coated overnight at 4°C with 5 µg/ml of the murine factor VIII light chain specific monoclonal antibody CLB-CAG 117 which was diluted in 50 mM NaHCO₃ (pH 9.5). Monoclonal antibody CLB-CAG 117 is directed against the C2-domain of factor VIII. The use of CLB-CAG 117 in the selection-protocol may result in elimination of recombinant phages that express immunoglobulin fragments directed against the C2-domain of factor VIII. Recombinant phages expressing the IgG4 specific immunoglobulin repertoire were prepared as described in Example 3. Recombinant phages were initially incubated on microtiter wells that contained immobilized CLB-CAG 117 for 2 hours in TBS supplemented with 3% HSA and 0.5 % Tween-20. The phage solution was removed and transferred to a second microtiter well (also coated with CLB-CAG 117) which had been preincubated with 1 µg/ml of factor VIII

light chain. Phages were allowed to bind to the immobilized factor VIII light chain for 2 hours at room temperature. Wells were washed extensively as described in Example 3 and bound phage were eluted with 100 mM triethylamine (pH 12). The eluted phage were neutralized by the addition of 1 M Tris-HCl pH 7.4 and the resulting solution was used to infect E. coli TG1 cells as described in Example 3.

Alternatively, purified factor VIII (5 µg/ml in 50 mM NaHCO₃ (pH 9.6)) was immobilized on immunotubes (Nunc, Life Technologies, Breda, The Netherlands). Recombinant phages diluted in TBS supplemented with 3% HSA were first incubated for 2 hours at room temperature in non-coated immunotubes. Subsequently, 1 ml of phage solution was removed and incubated for 2 hours at room temperature in Immunotubes coated with factor VIII. Following extensive washing (20 times with TBS/0.1% Tween-20 and 20 times with TBS) bound phage were eluted with 100 mM triethylamine (pH 12) and processed as outlined above.

The second, third and fourth round of panning were performed using the selection protocol described above. After the fourth round of panning 15 individual clones were picked and recombinant phage were tested for binding to the factor VIII light chain and the A3-C1 domain. Factor VIII light chain was purified as described previously. A construct expressing recombinant A3-C1 domain was prepared essentially as described previously (Sawamoto et al. 1998. Thrombosis and Haemostasis vol. 78, 62-68) and expressed in CHO-cells. The non-inhibitory murine monoclonal antibody CLB-CAG 12 directed against an epitope in the A3-C1 domain of factor VIII was immobilized on microtiter wells at a concentration of 5 µg/ml in 50 mM NaHCO₃ (pH 9.5). Wells were blocked for 1 hour in TBS supplemented with 3% HSA. Subsequently, wells were incubated with factor VIII light chain (1 µg/ml) or recombinant A3-C1 domain (0.06 nM) in 50 mM Tris HCl (pH 7.4), 1 M NaCl, 2% HSA for 2 hours at 37°C. Fifty µl of phage solution and an equal volume of TBS supplemented with 1% Tween-20 and 6% HSA were added to wells containing immobilized factor VIII light chain or A3-C1 domain. To monitor specific binding of the phage, wells that did not contain factor VIII light chain or recombinant A3-C1 domain were incubated with simultaneously with the phage solution. Phage were incubated at room temperature for 2 hours and were shaken at regular intervals. Wells were washed extensively with TBS supplemented with 0.1 % Tween-20. The presence of bound phage was

monitored as described in Example 3. The results of the analysis are depicted in Figure 8. In panel A, 15 clones selected in immunotubes that contain factor VIII are depicted. Of the 15 clones analyzed, 1 clone (clone 20) did not react with factor VIII light chain and recombinant A3-C1 domain suggesting that this clone does not encode an antibody fragment with specificity for the A3-C1 domain of factor VIII. Three clones (clone 17, 23 and 24) do react with the factor VIII light chain but fail to react with recombinant A3-C1 domain. Apparently, the epitope of these recombinant antibody fragments is localized in the C2-domain of factor VIII. The remaining 11 clones react both with the factor VIII light chain and the recombinant A3-C1 domain. In panel B, 15 clones selected in microtiter wells that contain CLB-CAG 117 and factor VIII light chain are depicted. Clones 31-45 all interact with the factor VIII light chain (black bars) and recombinant A3-C1 domain (hatched bars). This analysis shows that phage derived of clone 31-45 encode antibody fragments that bind specifically to the A3-C1 domain of factor VIII.

These results show that the protocol outlined above is suitable for the selection of recombinant antibody fragments that bind specifically to the A3-C1 domain of factor VIII. Using the methods disclosed in this example, it is feasible to isolate recombinant phage encoding antibody fragments specific for the A3-C1 domain from other patients with factor VIII inhibitors.

The nucleotide sequence of the variable heavy chain fragments of 26 clones that reacted specifically with recombinant A3-C1 domain was determined essentially as described in Example 4. The sequences obtained were aligned with heavy chain sequences in the database "V BASE" of the MRC Centre of Protein Engineering (Cambridge, UK). The 26 clones analyzed were encoded by four different VH-gene segments DP15, DP31 and DP49 and DP77. The amino acid sequence of the variable heavy chain fragments of clones B38, B18, B35 and B04 is listed in Figure 9A. The nucleotide sequence of these four clones is presented in Figures 9B-E.

EXAMPLE 9: Isolation and characteristics of anti-factor VIII antibodies that bind to the A2-domain of factor VIII.

An immunodominant region which constitutes a binding site for factor VIII inhibitors has been localized to the A2-domain of factor VIII (Healey et al. 1995, J. Biol. Chem. 270: 14505-14509). We characterized the anti-factor VIII antibodies in plasma of a patient with mild haemophilia A and an inhibitor,

essentially as outlined in example 1. Recombinant factor VIII fragments corresponding to the A2, A3-C1-C2 and C2-domain of factor VIII were metabolically labelled with [³⁵S]-methionine and used for the detection of anti-factor VIII antibodies in the patient's plasma. Reactivity with metabolically
5 labelled A2-domain and A3-C1-C2 domain was observed whereas only weak reactivity with metabolically labelled C2-domain was observed (data not shown). To determine the inhibitory capacity of both the anti-A2 and anti-A3-C1-C2 antibodies we performed neutralization experiments. The factor VIII inhibitor was diluted until a final value of 2 BU/ml and subsequently increasing
10 amounts of recombinant A2 or A3-C1-C2 were added. Addition of recombinant A2-domain resulted in almost complete neutralization of the factor VIII inhibitors present in patient's plasma. Addition of recombinant A3-C1-C2 only neutralized the factor VIII inhibitor to a limited extent (< 10%). These results show that the majority of factor VIII inhibitors are directed towards the A2-
15 domain of factor VIII. We assessed the subclass of the anti-factor VIII antibodies by enzyme linked sorbent assay. Both anti-A2 and anti-A3-C1-C2 antibodies consisted predominantly of subclass IgG4.

Peripheral blood lymphocytes of the patient were used to construct an IgG4-specific library as outlined in example 2. A library consisting of 1.9×10^6
20 clones was obtained. Recombinant phage expressing the IgG4-specific immunoglobulin repertoire of the patient were prepared as described in Example 3. Selection of phages binding to the A2-domain of factor VIII was performed by one of the following methods:

- 25 1. Purified factor VIII heavy chain (10 µg/ml) was immobilized on immunotubes (Nunc, Life Technologies, Breda, The Netherlands) in 50 mM NaHCO₃ (pH 9.5). Recombinant phages diluted in TBS supplemented with 3% HSA were first incubated for 2 hours at room temperature in non-coated immunotubes. Non-bound phage were transferred to an immunotube coated
30 with factor VIII. Following extensive washing (20 times with TBS/0.1% Tween-20 and 20 times with TBS) bound phage was eluted with 100 mM triethylamine (pH 12). Eluted phage was neutralized by the addition of 1 M Tris-HCl pH 7.4 and used to infect E.coli TG1 cells as described in Example 3.
- 35 2. Alternatively, the murine monoclonal antibody CLB-CAG 9, directed against amino acid sequence 713-740 in the A2-domain of factor VIII was

immobilized on microtiter wells at a concentration of 5 $\mu\text{g/ml}$ in 50 mM NaHCO_3 (pH 9.6). Purified factor VIII heavy chain (1 $\mu\text{g/ml}$) was then added and allowed to bind to CLB-CAG 9. Recombinant phage diluted in TBS 3% HSA and 0.5% Tween 20 were first incubated in microtiter wells containing
5 only immobilized CLB-CAG 9. After 2 hours non-bound phage were transferred to a microtiter well which contained immobilized factor VIII heavy chain. Phage were allowed to bind to the factor VIII heavy chain for 2 hours at room temperature. Wells were washed extensively (see above) and finally bound phage were eluted with 100 mM triethylamine (pH 12) and processed as
10 outlined above.

After four rounds of selection individual clones were picked and binding of recombinant phage to factor VIII heavy chain was evaluated by an enzyme linked immuno sorbent assay. Monoclonal antibody ESH5 (American
15 Diagnostica, Greenwich, CT, USA) was immobilized onto microtiter wells at a concentration of 5 $\mu\text{g/ml}$ in 50 mM NaHCO_3 (pH 9.6). Purified factor VIII heavy chain (1 $\mu\text{g/ml}$) was added and incubated for 2 hours at 37°C. Subsequently, recombinant phage, diluted 1 to 1 in 50 mM Tris-HCl pH 7.4, 1 M NaCl and 2% HSA, was added and incubated for 2 hours at room
20 temperature. The amount of recombinant phage bound was determined as described in Example 3. The results of the analysis is given in Figure 10. Twenty clones which were selected in microtiter wells in which factor VIII heavy chain was immobilized by CLB-CAG 9 were analyzed. Eleven out of 20 clones bound specifically to the heavy chain of factor VIII
25 (Figure 10A). Clones that were selected by immobilized factor VIII heavy chain in immunotubes were also analyzed. Fifteen out of 20 clones bound specifically to the factor VIII heavy chain (Figure 10B). These results show that the protocol outlined above permits the isolation of anti-factor VIII antibodies that are directed against the heavy chain (A1-A2) of factor VIII. Using the
30 methods disclosed in this example it is feasible to isolate anti-factor VIII antibodies from the repertoire of additional patients with factor VIII inhibitors directed against the A2-domain.

The nucleotide sequence of the variable heavy chain fragments of 26 clones that reacted specifically with the factor VIII heavy chain were
35 determined essentially as described in Example 4. The sequences obtained were aligned with heavy chain sequences in the database "V BASE" of the MRC

Centre of Protein Engineering (Cambridge, UK). The 26 clones analyzed were encoded by two different VH-gene segments DP10 and DP47 (Figure 11A). The nucleotide sequence of the variable heavy chain of these clones is listed in Figure 11B and C.

5

EXAMPLE 10: Factor VIII specific recombinant antibody fragments can neutralize the activity of factor VIII inhibitors present in plasma of patients with haemophilia.

In example 7, we have shown that scFv-EL14 and scFv-IT2 neutralize the inhibitory activity of the murine monoclonal antibody CLB-CAG 117. We tested whether scFv-EL14 can also neutralize factor VIII inhibitors present in plasma of haemophilia A patients. First, we tested plasma of the patient with acquired haemophilia from whom the recombinant antibody fragments were derived. As described in example 1, recombinant C2-domain was capable of neutralizing 20% of the factor VIII inhibitor in patient's plasma (Table III). The effect of scFv-EL14 was evaluated in a similar set-up. Plasma was diluted till a final value of 2 BU/ml and increasing amounts of scFv-EL14 were added. ScFv-EL14 could neutralize about 20% of the total activity of factor VIII inhibitor in patient's plasma. These results suggest that scFv directed against the C2-domain prevent binding of factor VIII inhibitory antibodies that bind to the C2-domain of factor VIII.

Next, we tested two plasma samples derived of patients with congenital haemophilia A and factor VIII inhibitors. The relative contribution of the C2-domain to the total amount of factor VIII inhibitor for both samples ranged between 40% and 90%. Neutralization experiments indicate that addition of increasing concentrations of scFv-EL14 results in significant reduction of the levels of factor VIII inhibitor in plasma of these two patients with congenital haemophilia A. These results confirm that scFv-EL14 alleviates binding of human factor VIII inhibitors to the C2-domain. Our findings show that scFv-EL14 shields antigenic sites that are present in the C2-domain of factor VIII. This property of scFv-EL14 can be utilized to prevent binding of factor VIII inhibitors to the C2-domain of administered factor VIII in haemophilia A patients with an inhibitor. It has been firmly established that factor VIII inhibitors often recognize multiple epitopes that have been localized to the A2-, A3- and C2 domain of factor VIII. In this example the neutralizing activity of the C2-domain specific scFv-EL14 on the biological activity of factor VIII

inhibitors is described.

In examples 8 and 9 we have disclosed methods to obtain recombinant antibodies that specifically react with the A3-C1 domain and heavy chain (A1-A2) of factor VIII. In this example, we have shown that recombinant antibodies
5 directed against the C2-domain of factor VIII can shield antigenic sites on factor VIII. Similarly, anti-A3-C1 antibodies and anti-A2-antibodies described in examples 8 and 9 can be tested for their ability to compete with factor VIII inhibitors for binding to factor VIII. The anti-A3-C1 and anti-A1-A2 antibodies disclosed in this invention may also be used for treatment of patients with
10 inhibitors, which react with the A3-C1 and/or the A1-A2-domain.

Table I: Nucleotide sequences of clones expressing recombinant antibodies with specificity for the factor VIII light chain. Based on the nucleotide sequence 55 of the 60 clones analyzed could be arranged as depicted below. In the first column clones with the same nucleotide sequence are arranged in four groups. The number of clones corresponding to this group is given in brackets. Clone EL5, EL25 and IT2 are related as indicated in Figure 4B. In the second column the heavy chain family to which these clones belong is depicted. All clones analyzed belong to the VH1-family. In the third column the germline segment is depicted. Clone EL5, EL25 and IT2 belong to germline segment DP-14 while clone EL14 belongs to germline sequence DP-10. In the fourth column the number of mutations in the different clones is depicted. The first number corresponds to the number of nucleotide mutations while the second one corresponds to the number of amino acid changes. The sequences were compared with the nucleotide and amino acid sequences of the germline segments indicated in the Table.

Clone	VH family	Germline segment	Mutations
EL5 (3)	VH1	DP-14	20/12
EL14 (14)	VH1	DP-10	18/12
EL25 (5)	VH1	DP-14	19/11
IT2 (33)	VH1	DP-14	20/13

PCT pat. appln. no. PCT/NL99/00285
Our letter of

Claims

1. A polynucleotide in substantially isolated form, comprising a contiguous nucleotide sequence (a) coding for a human antibody with factor VIII specificity, or (b) complementary to a nucleotide sequence coding for a human antibody with factor VIII specificity, or (c) capable of selectively hybridizing under stringent conditions to nucleotide sequence (a) or (b).
2. A polynucleotide according to claim 1, wherein said contiguous nucleotide sequence is at least 8, preferably at least 10 nucleotides.
3. A probe or primer which comprises a polynucleotide according to claim 1 or claim 2, optionally further comprising a detectable label, such as a radioactive atom or group, an enzyme, a fluorescent or luminescent group, a dye or biotin.
4. An assay kit for detecting nucleic acid coding for a human antibody with factor VIII specificity, comprising a probe or primer according to claim 3 in a suitable container.
5. A nucleic acid amplification and detection kit for detecting nucleic acid coding for a human antibody with factor VIII specificity, comprising a pair of primers according to claim 3 capable of priming the synthesis of cDNA, and optionally further comprising a probe according to claim 3 capable of selectively hybridizing to (the complement of) a region of the nucleic acid to be detected between and not including the sequences from which the primers are derived.
6. A method for assaying a sample for the presence or absence of nucleic acid coding for a human antibody with factor VIII specificity, comprising contacting the sample with a probe according to claim 3 under conditions that allow the selective hybridization of said probe to the (complement of the) nucleic acid to be detected in the sample, and determining whether polynucleotide duplexes comprising said probe are formed.
7. A method for assaying a sample for the presence or absence of nucleic acid coding for a human antibody with factor VIII specificity, comprising subjecting nucleic acid present in the sample to a nucleic acid amplification process using a

pair of primers according to claim 3 capable of priming the synthesis of cDNA, contacting the nucleic acid resulting from the amplification process with a probe according to claim 3 under conditions that allow the selective hybridization of said probe to the (complement of the) nucleic acid to be detected in the sample, and determining whether polynucleotide duplexes comprising said probe are formed.

8. A method of producing a recombinant polypeptide, comprising providing a polynucleotide coding for said polypeptide, preparing a recombinant vector containing said polynucleotide operably linked to a control sequence capable of providing for the expression of the polynucleotide by a host cell, transforming a host cell with said recombinant vector, growing said host cell under conditions that provide for the expression of the polynucleotide and optionally isolating the thus produced polypeptide, wherein said polynucleotide codes for a human antibody with factor VIII specificity, or a fragment or derivative thereof capable of specific binding to factor VIII.

9. A polypeptide in substantially isolated form, comprising a contiguous amino acid sequence corresponding to or mimicking a fragment or derivative of a human antibody with factor VIII specificity capable of specific binding to factor VIII.

10. A polypeptide according to claim 9, wherein said contiguous amino acid sequence is capable of reducing the activity of factor VIII inhibiting antibodies.

11. A polypeptide according to claim 9 or claim 10, wherein said fragment is (part of) a variable region of the heavy chain or light chain of said antibody.

12. A polypeptide according to claim 9 or claim 10, wherein said derivative is a single chain Fv fragment of said antibody.

13. An antibody in substantially isolated form, comprising a recombinant human antibody with factor VIII specificity or an anti-idiotypic antibody directed against a human antibody with factor VIII specificity.

14. A pharmaceutical composition for the treatment of factor VIII inhibition in a human individual, comprising a polypeptide according to any one of claims 9-12 or an antibody according to claim 13, together with a pharmaceutically acceptable carrier.

15. A composition according to claim 14, which further contains factor VIII or a substitute of factor VIII.

16. A method of treatment of factor VIII inhibition in a human individual, comprising administering to said individual a polypeptide according to any one of claims 9-12 or an antibody according to claim 13, optionally together with factor VIII or a substitute of factor VIII.

17. A polypeptide capable of specific binding to factor VIII and interference with the activity of factor VIII inhibitors, which polypeptide comprises the variable part of the heavy chain of a human antibody with factor VIII specificity or a part thereof which at least includes the CDR3 region.

- 5 18. A polypeptide according to claim 17 which essentially consists of (a) the CDR3 region of the variable part of the heavy chain of a human antibody with factor VIII specificity, (b) an antibody fragment containing the variable part of the heavy chain of a human antibody with factor VIII specificity, or (c) a single chain Fv fragment containing the variable part of the heavy chain of a human antibody
10 with factor VIII specificity.

19. A polynucleotide in substantially isolated form, coding for a polypeptide according to claim 17 or 18.

- 15 20. A pharmaceutical composition for the treatment of factor VIII inhibition in a human individual, comprising a polypeptide according to claim 17 or 18 together with a pharmaceutically acceptable carrier.

21. A pharmaceutical composition according to claim 20, which further contains factor VIII or a substitute of factor VIII.

- 20 22. A method of treatment of factor VIII inhibition in a human individual, comprising administering to said individual a polypeptide according to claim 17 or 18.

23. A method of treatment of factor VIII inhibition in a human individual, comprising administering to said individual a polypeptide according to claim 17 or 18 together with factor VIII or a substitute of factor VIII.

25

1/20

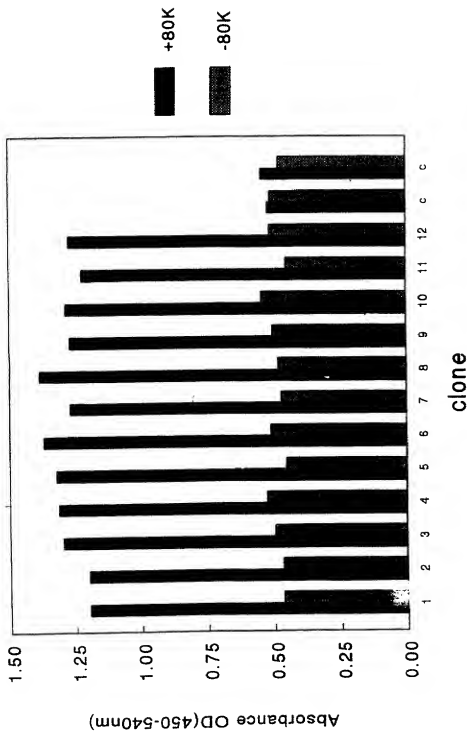


Figure 1

2/20

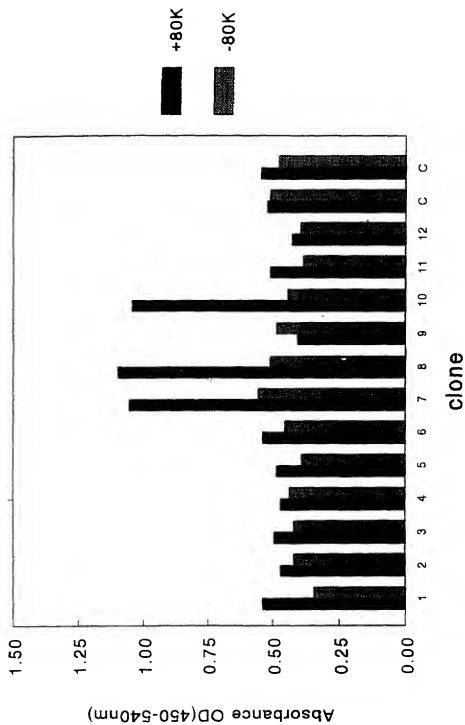


Figure 2

3/20

Figure 3

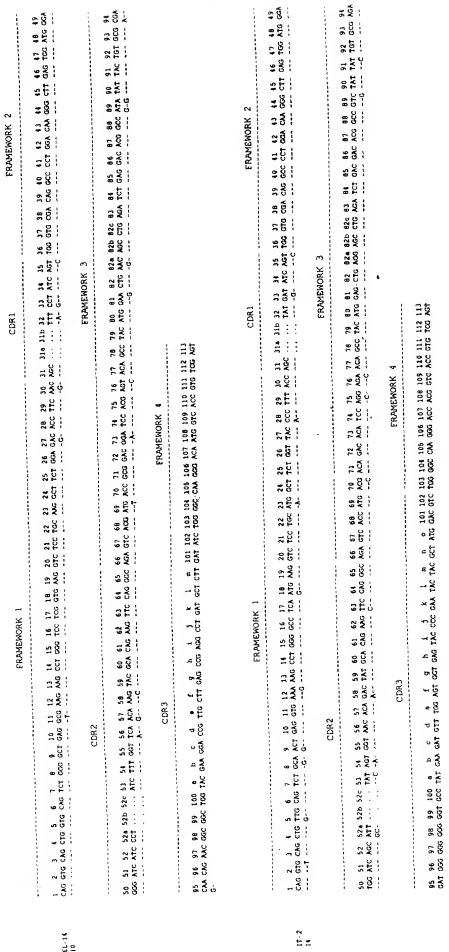


Figure 4A

SUBSTITUTE SHEET (RULE 26)

	FR1	FR2	FR3	FR4
1	1	1	1	1
2	2	2	2	2
3	3	3	3	3
4	4	4	4	4
5	5	5	5	5
6	6	6	6	6
7	7	7	7	7
8	8	8	8	8
9	9	9	9	9
10	10	10	10	10
11	11	11	11	11
12	12	12	12	12
13	13	13	13	13
14	14	14	14	14
15	15	15	15	15
16	16	16	16	16
17	17	17	17	17
18	18	18	18	18
19	19	19	19	19
20	20	20	20	20
21	21	21	21	21
22	22	22	22	22
23	23	23	23	23
24	24	24	24	24
25	25	25	25	25
26	26	26	26	26
27	27	27	27	27
28	28	28	28	28
29	29	29	29	29
30	30	30	30	30
31	31	31	31	31
32	32	32	32	32
33	33	33	33	33
34	34	34	34	34
35	35	35	35	35
36	36	36	36	36
37	37	37	37	37
38	38	38	38	38
39	39	39	39	39
40	40	40	40	40
41	41	41	41	41
42	42	42	42	42
43	43	43	43	43
44	44	44	44	44
45	45	45	45	45
46	46	46	46	46
47	47	47	47	47
48	48	48	48	48
49	49	49	49	49
50	50	50	50	50
51	51	51	51	51
52	52	52	52	52
53	53	53	53	53
54	54	54	54	54
55	55	55	55	55
56	56	56	56	56
57	57	57	57	57
58	58	58	58	58
59	59	59	59	59
60	60	60	60	60
61	61	61	61	61
62	62	62	62	62
63	63	63	63	63
64	64	64	64	64
65	65	65	65	65
66	66	66	66	66
67	67	67	67	67
68	68	68	68	68
69	69	69	69	69
70	70	70	70	70
71	71	71	71	71
72	72	72	72	72
73	73	73	73	73
74	74	74	74	74
75	75	75	75	75
76	76	76	76	76
77	77	77	77	77
78	78	78	78	78
79	79	79	79	79
80	80	80	80	80
81	81	81	81	81
82	82	82	82	82
83	83	83	83	83
84	84	84	84	84
85	85	85	85	85
86	86	86	86	86
87	87	87	87	87
88	88	88	88	88
89	89	89	89	89
90	90	90	90	90
91	91	91	91	91
92	92	92	92	92
93	93	93	93	93
94	94	94	94	94
95	95	95	95	95
96	96	96	96	96
97	97	97	97	97
98	98	98	98	98
99	99	99	99	99
100	100	100		

Figure 4B

6/20

CDR3

EL-14	QQNGGWYEGPLLEPRPD--ALDI
	. .
IT-2	DGGGGAYEDVWSGEYPEYYAMDV

Figure 4C

7/20

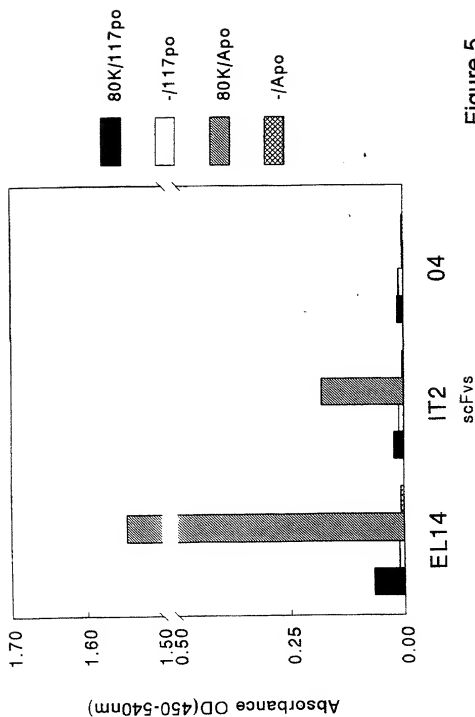


Figure 5

8/20

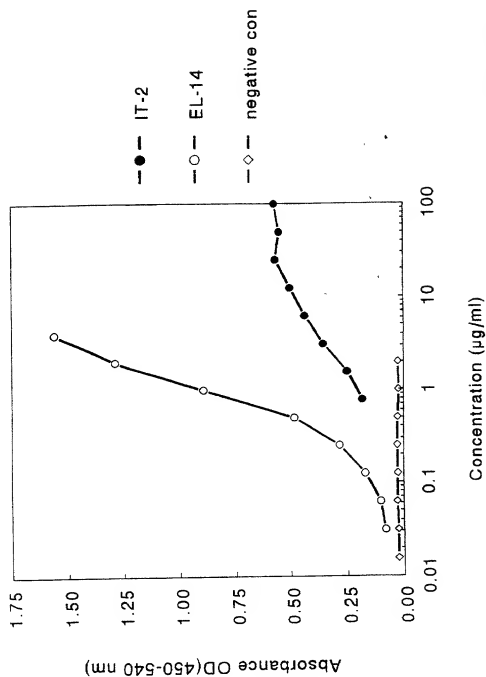


Figure 6

9/20

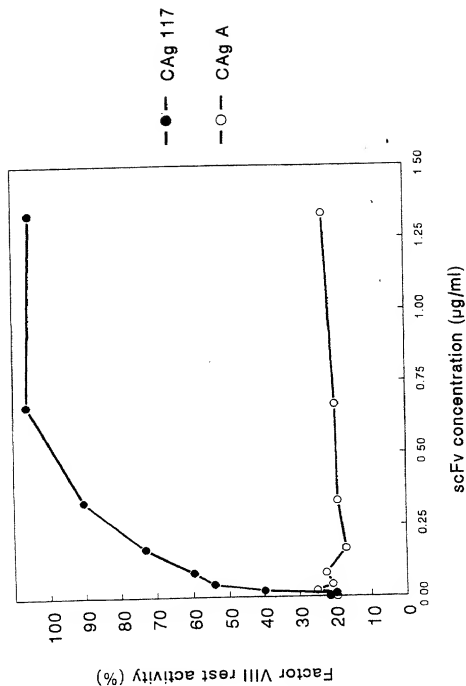


Figure 7A NEUTRALIZATION BY scFv EL-14

10/20

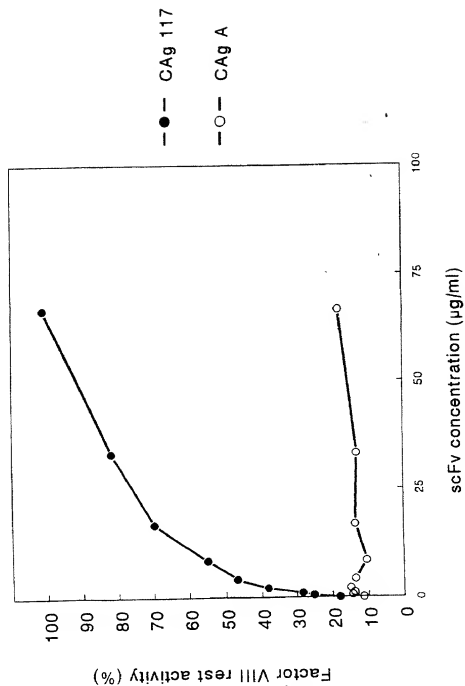
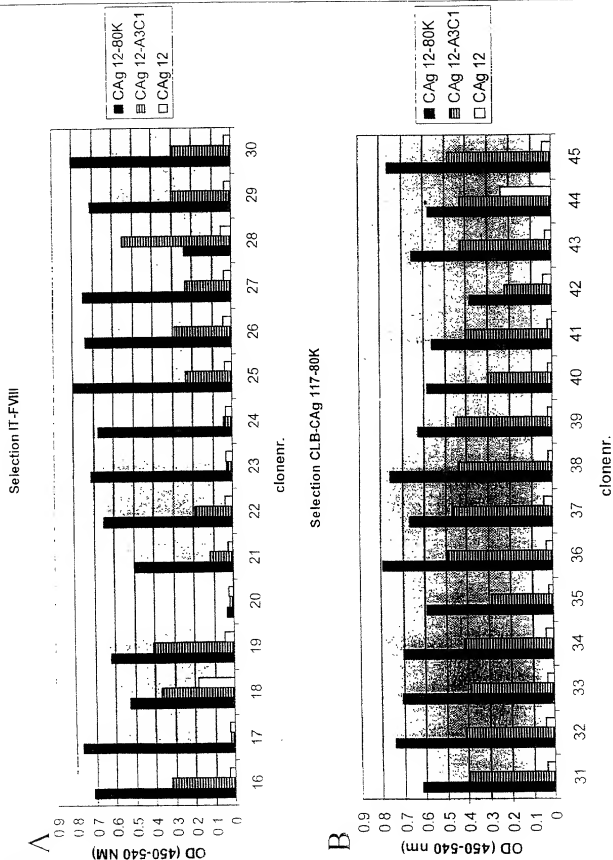


Figure 7B NEUTRALIZATION BY scFv IT-2

Figure 8



12/20

Deduced protein sequences of isolated FVIII A3-C1 specific scFv

[illegible]

Figure 9A

13/20

-1 -Gln-Val-Gln-Leu-Leu-Gln-Ser-Ala-Ala-Asp-Val-Lys-Lys-Pro-Gly-Ala-Ser-
 CAGGTGCAGC TGTTGCAGTC TGCAGCTGAC GTGAAGAAGC CTGGGGCCTC 50
 GTCCACGTCG ACAACGTCAG ACGTCGACTG CACTTCTTCG GACCCCGGAG

-1 -Val-Lys-Val-Ser-Cys-Thr-Ala-Ser-Gly-Tyr-Ile-Phe-Thr-Ser-Tyr-Asp-Ile-
 AGTGAAGGTC TCCTGTACGG CTTCTGGATA CATCTTCACC AGTTATGATA 100
 TCACTTCCAG AGGACATGCC GAAGACCTAT GTAGAAGTGG TCAATACTAT

-1 -Asn-Trp-Val-Arg-Gln-Ala-Thr-Gly-Gln-Gly-Leu-Glu-Trp-Met-Gly-Trp-
 TCAACTGGGT GCGACAGGCC ACTGGACAAG GGCTTGAGTG GATGGGATGG 150
 AGTTGACCCA CGCTGTCCGG TGACCTGTTC CCGAACTCAC CTACCTACC

-1 Met-Asn-Pro-Asn-Ser-Gly-Asn-Ala-Gly-Phe-Ala-Gln-Lys-Phe-Lys-Gly-Arg-
 ATGAATCCTA ACAGTGGTAA CGCAGGCTTT GCACAGAAGT TTAAGGGCAG 200
 TACTTAGGAT TGTCACCATT GCGTCCGAAA CGTGTCTTCA AATTCCCGTC

-1 -Leu-Thr-Leu-Thr-Arg-Asp-Thr-Ser-Thr-Ser-Thr-Ala-Tyr-Met-Glu-Leu-Arg
 ACTCACCTTG ACCAGGGACA CTTCCACAAG CACAGCCTAC ATGGAGCTGA 250
 TGAGTGGAAC TGGTCCCTGT GAAGGTGTTC GTGTCGGATG TACCTCGACT

-1 -Arg-Leu-Glu-Ser-Glu-Asp-Thr-Ala-Val-Tyr-Tyr-Cys-Ala-Arg-Cys-Asp-
 GGAGACTGGA ATCTGAGGAC ACGGCCGTGT ATTACTGTGC GAGATGTGAC 300
 CCTCTGACCT TAGACTCCTG TGCCGGCACA TAATGACACG CTCTACACTG

-1 -Thr-Thr-Leu-Leu-Ile-Trp-Phe-Gly-Pro-Ala-Pro-Tyr-Tyr-Asp-Ser-Trp-Gly-
 ACCACACTCT TAATCTGGTT CGGGCCCGCC CCCTACTATG ACTCCTGGGG 350
 TGGTGTGAGA ATTAGACCAA GCCCGGGCGG GGGATGATAC TGAGGACCCC

-1 -Gln-Gly-Thr-Leu-Val-
 CCAGGGAAct CTAGTC 400
 GGTCCCTTGA GATCAG

Figure 9B

14/20

+1 -Gln-Val-Gln-Leu-Val-Gln-Ser-Gly-Gly-Gly-Leu-Val-Gln-Pro-Gly-Lys-Ser-
 CAGGTGCAAC TGGTGCAGTC TGGGGGAGGC TTGGTACAGC CTGGCAAGTC 50
 GCGACGTTG ACCACGTCAG ACCCCCTCCG AACCATGTCG GACCGTTCAG

+1 -Leu-Arg-Leu-Ser-Cys-Ala-Ala-Ser-Gly-Phe-Thr-Phe-Gly-Asp-Tyr-Ala-Ile-
 CCTGAGACTC TCCTGTGCAG CCTCTGGATT CACATTGGC GATTATGCCA 100
 GGACTCTGAG AGGACACGTC GGAGACCTAA GTGTAAACCG CTAATACGGT

+1 -His-Trp-Val-Arg-Gln-Ala-Pro-Gly-Glu-Gly-Leu-Glu-Trp-Val-Ser-Gly-
 TACACTGGGT CCGGCAAGCT CCAGGGGAGG GCCTGGAGTG GGTCTCAGGT 150
 ATGTGACCCA GGCCGTTCTG GGTCCCTCC CGGACCTCAC CCAGAGTCCA

+1 -Val-Thr-Trp-Ser-Gly-Thr-Thr-Ile-Gly-Phe-Ala-Asp-Ser-Val-Lys-Gly-Arg-
 GTTACTTGGA GTGGTACTAC TATAGGCTTT GCGGACTCTG TGAAGGGCCG 200
 CAATGAACCT CACCATGATG ATATCCGAAA CGCCTGAGAC ACTTCCCGGC

+1 -Phe-Thr-Ile-Ser-Arg-Asp-Asn-Ala-Lys-Asn-Ser-Leu-Tyr-Leu-Tyr-Met-Asn-
 ATTCAACATC TCCAGAGACA ACGCCAAGAA TTCCCTGTAT CTGTACATGA 250
 TAAGTGGTAG AGGTCTCTGT TCGGTTCTT AAGGGACATA GACATGTACT

+1 -Ser-Leu-Arg-Ala-Glu-Asp-Thr-Ala-Leu-Tyr-Tyr-Cys-Ala-Leu-Pro-Tyr-
 ACAGTCTGAG AGCTGAAGAC ACGGCCTTGT ATTATTGTGC CTTACCATAT 300
 TGTCAGACTC TCGACTTCTG TGCCGGAACA TAATAACACG GAATGGTATA

+1 -Ile-Asn-Ser-Ser-Asn-Tyr-Arg-Arg-Gly-Val-Ala-Ala-Phe-Asp-Ile-Trp-Gly-
 ATCAACTCGT CCAACTACAG AAGAGGGGTC GCTGCCTTCG ATATCTGGGG 350
 TAGTTGAGCA GGTGTATGTC TTCTCCCCAG CGACGGAAGC TATAGACCCC

+1 -Gln-Gly-Thr-Met-Val-Thr-Val-Ser-
 CCAAGGGACA ATGGTCACCG TGTCGAGT 400
 GGTTCCCTGT TACCACTGGC ACAGCTCA

Figure 9C

15/20

+1 Glu Val Gln Leu Val Glu Ser Gln Gly Gly Leu Val Gln Pro Gly Arg Ser
 GAGGTGCAGC TGGTGGAGTC TGGGGGAGGC TTGGTACAGC CTGGGAGGTC 50
 CTCCACGTCG ACCACCTCAG ACCCCCTCCG AACCATGTG GACCTCCAG

+1 Leu Arg Leu Ser Cys Val Asp Ser Gly Leu Thr Phe Ser Ser Tyr Gly Met
 CCTGAGACTC TCCTGTGTAG ACTCTGGACT CACCTTCAGT AGTTATGGCA 100
 GGACTCTGAG AGGACACATC TGAGACCTGA GTGGAAGTCA TCAATACCGT

+1 His Trp Val Arg Gln Ala Pro Gly Ala Gly Leu Glu Trp Val Ala Val
 TGCCTGGGT CCGCCAGGCT CCAGGCGCGG GGCTGGAGTG GGTGGCCGTT 150
 ACGTGACCCA GGCGTCCGA GGTCCGCGCC CCGACCTCAC CCACCGCAA

- Ile Ser Tyr Asp Gly Asn Asp Lys Tyr Tyr Ala Asp Ser Val Lys Gly Arg
 ATTCATACG ACGGAAATGA TAAATATTAT GCAGACTCCG TGAAGGGCCG 200
 TAAAGTATGC TGCCTTTACT ATTTATAATA CGTCTGAGGC ACTTCCCGGC

+1 Phe Ala Ile Ser Arg Asp Asn Ala Lys Asn Thr Leu Tyr Leu Gln Met Asn
 ATTCGCCATC TCCAGAGACA ATGCCAAGAA CACGCTGTAT CTGCAAATGA 250
 TAAGCGGTAG AGGTCTCTGT TACGTTTCTT GTGCGACATA GACGTTTACT

+1 Ser Leu Thr Ile Gln Asp Thr Ala Val Tyr Tyr Cys Ala Lys Asp Leu
 ACAGCCTGAC AATAGAGGAC ACGGCTGTCT ATTATTGTGC GAAAGATCTC 300
 TGTCGGACTG TTATCTCCTG TGCCGACAGA TAATAACACG CTTTCTAGAG

+1 Ile Glu Ser Asn Ile Ala Glu Ala Leu Trp Gly Gln Gly Thr Leu Val Thr
 ATAGAATCAA ATATTGCGGA GGCCCTCTGG GGCCAGGGAA CCCTGGTCAC 350
 TATCTTAGTT TATAACGCCT CCGGGAGACC CCGTCCCTT GGGACCAAGT

- Val Ser Ser
 CGTGTGAGT
 GCACAGCTCA 400

Figure 9D

16/20

+1 : Glu₁ Val₁ Glu₁ Leu₁ Val₁ Lys₁ Ser₁ Gly₁ Glu₁ Gly₁ Leu₁ Val₁ Lys₁ Pro₁ Gly₁ Gly₁ Ser₁
 GAGGTGCAGC TGGTGAAGTC TGGGGAAGGC CTGGTCAAGC CTGGGGGGTC 50
 CTCCACGTCG ACCACTTCAG ACCCCTTCCG GACCAGTTCCG GACCCCCCAG

-1 : Leu₁ Arg₁ Leu₁ Ser₁ Cys₁ Ala₁ Ala₁ Ser₁ Gly₁ Phe₁ Thr₁ Phe₁ Arg₁ Arg₁ Tyr₁ Asp₁ Ile₁
 CCTGAGACTC TCCTGTGCAG CCTCTGGATT CACCTTCAGG AGATATGATA 100
 GGACTCTGAG AGGACACGTC GGAGACCTAA GTGGAAGTCC TCTATACTAT

+1 : His₁ Trp₁ Val₁ Arg₁ Glu₁ Thr₁ Pro₁ Gly₁ Lys₁ Gly₁ Leu₁ Glu₁ Trp₁ Val₁ Ser₁ Ser₁
 TCCACTGGGT CCGCCAGACT CCAGGGAAGG GCCTGGAGTG GGTCTCATCC 150
 AGGTGACCCA GCGGGTCTGA GGTCCCTTCC CGGACCTCAC CCAGAGTAGG

+1 : Ile₁ Ser₁ Ser₁ Gly₁ Gly₁ Asn₁ Tyr₁ Ile₁ Asp₁ Tyr₁ Ala₁ Asp₁ Ser₁ Val₁ Lys₁ Gly₁ Arg₁
 ATCAGTAGTG GTGGTAATTA CATAGACTAC GCAGACTCTG TGAAGGGCCG 200
 TAGTCATCAC CACCATTAAAT GTATCTGATG CGTCTGAGAC ACTTCCCGGC

+1 : Phe₁ Thr₁ Ile₁ Ser₁ Arg₁ Asp₁ Asn₁ Ala₁ Asn₁ Asn₁ Val₁ Val₁ Tyr₁ Leu₁ Glu₁ Met₁ Asn₁
 ATTCAACATC TCCAGAGACA ACGCCAACAA TGTGTCTTAT CTACAATGA 250
 TAAAGTGGTAG AGGTCTCTGT TGCGTTGTGT ACAACAGATA GATGTTTACT

+1 : Ser₁ Leu₁ Arg₁ Ala₁ Glu₁ Asn₁ Met₁ Ala₁ Val₁ Tyr₁ Phe₁ Cys₁ Ala₁ Arg₁ Asp₁ Gly₁
 ACAGCCTGAG AGCCGAGGAC ATGGCTGTCT ATTTCTGTGC GAGAGATGGG 300
 TGTCGGACTC TCGGCTCCTG TACCGACAGA TAAAGACACG CTCTCTACCC

+1 : Thr₁ Ile₁ Phe₁ Gly₁ Ser₁ Ala₁ Ala₁ Thr₁ Trp₁ Arg₁ Ala₁ Phe₁ Asp₁ Ile₁ Trp₁ Gly₁ Arg₁
 ACCATTTTGG GATCGGCGGC GACCTGGCGG GCTTTTGATA TCTGGGGCCG 350
 TGCTAAAAAC CTAGCCGCCG CTGGACCGCC CGAAACTAT AGACCCCGCC

-1 : Gly₁ Thr₁ Met₁ Val₁ Thr₁ Val₁ Ser₁ Ser₁
 GGGGACAATG GTCACCGTGT CGAGT 400
 CCCCTGTTAC CAGTGGCACA GCTCA

Figure 9E

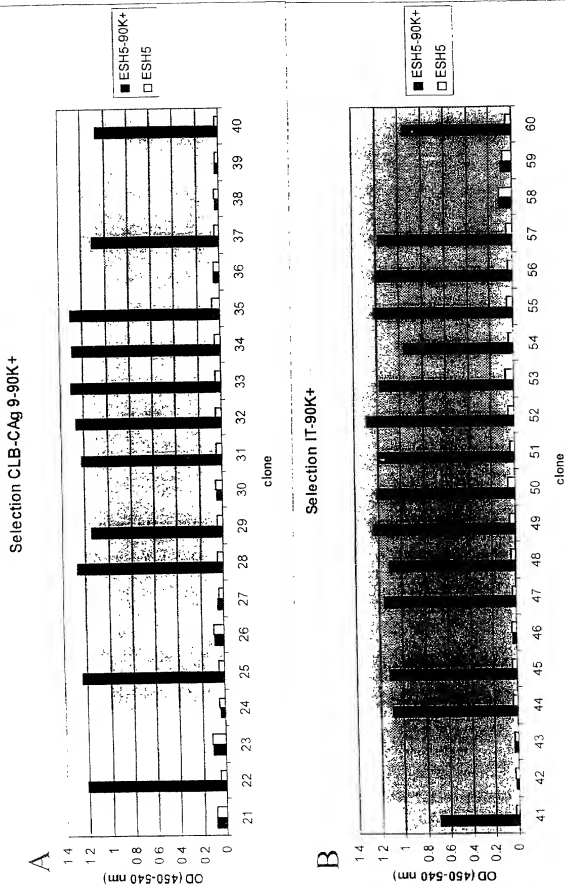


Figure 10

18/20

Deduced protein sequences of isolated FVIII A2 specific scFv

	FR1	FR2	FR3	FR4	FR5	FR6	FR7	FR8	FR9	FR10	FR11	FR12	FR13	FR14	FR15	FR16	FR17	FR18	FR19	FR20	FR21	FR22	FR23	FR24	FR25	FR26	FR27	FR28	FR29	FR30	FR31	FR32	FR33	FR34	FR35	FR36	FR37	FR38	FR39	FR40	FR41	FR42	FR43	FR44	FR45	FR46	FR47	FR48	FR49	FR50	FR51	FR52	FR53	FR54	FR55	FR56	FR57	FR58	FR59	FR60	FR61	FR62	FR63	FR64	FR65	FR66	FR67	FR68	FR69	FR70	FR71	FR72	FR73	FR74	FR75	FR76	FR77	FR78	FR79	FR80	FR81	FR82	FR83	FR84	FR85	FR86	FR87	FR88	FR89	FR90	FR91	FR92	FR93	FR94	FR95	FR96	FR97	FR98	FR99	FR100	FR101	FR102	FR103	FR104	FR105	FR106	FR107	FR108	FR109	FR110	FR111	FR112	FR113	FR114	FR115	FR116	FR117	FR118	FR119	FR120	FR121	FR122	FR123	FR124	FR125	FR126	FR127	FR128	FR129	FR130	FR131	FR132	FR133	FR134	FR135	FR136	FR137	FR138	FR139	FR140	FR141	FR142	FR143	FR144	FR145	FR146	FR147	FR148	FR149	FR150	FR151	FR152	FR153	FR154	FR155	FR156	FR157	FR158	FR159	FR160	FR161	FR162	FR163	FR164	FR165	FR166	FR167	FR168	FR169	FR170	FR171	FR172	FR173	FR174	FR175	FR176	FR177	FR178	FR179	FR180	FR181	FR182	FR183	FR184	FR185	FR186	FR187	FR188	FR189	FR190	FR191	FR192	FR193	FR194	FR195	FR196	FR197	FR198	FR199	FR200	FR201	FR202	FR203	FR204	FR205	FR206	FR207	FR208	FR209	FR210	FR211	FR212	FR213	FR214	FR215	FR216	FR217	FR218	FR219	FR220	FR221	FR222	FR223	FR224	FR225	FR226	FR227	FR228	FR229	FR230	FR231	FR232	FR233	FR234	FR235	FR236	FR237	FR238	FR239	FR240	FR241	FR242	FR243	FR244	FR245	FR246	FR247	FR248	FR249	FR250	FR251	FR252	FR253	FR254	FR255	FR256	FR257	FR258	FR259	FR260	FR261	FR262	FR263	FR264	FR265	FR266	FR267	FR268	FR269	FR270	FR271	FR272	FR273	FR274	FR275	FR276	FR277	FR278	FR279	FR280	FR281	FR282	FR283	FR284	FR285	FR286	FR287	FR288	FR289	FR290	FR291	FR292	FR293	FR294	FR295	FR296	FR297	FR298	FR299	FR300	FR301	FR302	FR303	FR304	FR305	FR306	FR307	FR308	FR309	FR310	FR311	FR312	FR313	FR314	FR315	FR316	FR317	FR318	FR319	FR320	FR321	FR322	FR323	FR324	FR325	FR326	FR327	FR328	FR329	FR330	FR331	FR332	FR333	FR334	FR335	FR336	FR337	FR338	FR339	FR340	FR341	FR342	FR343	FR344	FR345	FR346	FR347	FR348	FR349	FR350	FR351	FR352	FR353	FR354	FR355	FR356	FR357	FR358	FR359	FR360	FR361	FR362	FR363	FR364	FR365	FR366	FR367	FR368	FR369	FR370	FR371	FR372	FR373	FR374	FR375	FR376	FR377	FR378	FR379	FR380	FR381	FR382	FR383	FR384	FR385	FR386	FR387	FR388	FR389	FR390	FR391	FR392	FR393	FR394	FR395	FR396	FR397	FR398	FR399	FR400	FR401	FR402	FR403	FR404	FR405	FR406	FR407	FR408	FR409	FR410	FR411	FR412	FR413	FR414	FR415	FR416	FR417	FR418	FR419	FR420	FR421	FR422	FR423	FR424	FR425	FR426	FR427	FR428	FR429	FR430	FR431	FR432	FR433	FR434	FR435	FR436	FR437	FR438	FR439	FR440	FR441	FR442	FR443	FR444	FR445	FR446	FR447	FR448	FR449	FR450	FR451	FR452	FR453	FR454	FR455	FR456	FR457	FR458	FR459	FR460	FR461	FR462	FR463	FR464	FR465	FR466	FR467	FR468	FR469	FR470	FR471	FR472	FR473	FR474	FR475	FR476	FR477	FR478	FR479	FR480	FR481	FR482	FR483	FR484	FR485	FR486	FR487	FR488	FR489	FR490	FR491	FR492	FR493	FR494	FR495	FR496	FR497	FR498	FR499	FR500	FR501	FR502	FR503	FR504	FR505	FR506	FR507	FR508	FR509	FR510	FR511	FR512	FR513	FR514	FR515	FR516	FR517	FR518	FR519	FR520	FR521	FR522	FR523	FR524	FR525	FR526	FR527	FR528	FR529	FR530	FR531	FR532	FR533	FR534	FR535	FR536	FR537	FR538	FR539	FR540	FR541	FR542	FR543	FR544	FR545	FR546	FR547	FR548	FR549	FR550	FR551	FR552	FR553	FR554	FR555	FR556	FR557	FR558	FR559	FR560	FR561	FR562	FR563	FR564	FR565	FR566	FR567	FR568	FR569	FR570	FR571	FR572	FR573	FR574	FR575	FR576	FR577	FR578	FR579	FR580	FR581	FR582	FR583	FR584	FR585	FR586	FR587	FR588	FR589	FR590	FR591	FR592	FR593	FR594	FR595	FR596	FR597	FR598	FR599	FR600	FR601	FR602	FR603	FR604	FR605	FR606	FR607	FR608	FR609	FR610	FR611	FR612	FR613	FR614	FR615	FR616	FR617	FR618	FR619	FR620	FR621	FR622	FR623	FR624	FR625	FR626	FR627	FR628	FR629	FR630	FR631	FR632	FR633	FR634	FR635	FR636	FR637	FR638	FR639	FR640	FR641	FR642	FR643	FR644	FR645	FR646	FR647	FR648	FR649	FR650	FR651	FR652	FR653	FR654	FR655	FR656	FR657	FR658	FR659	FR660	FR661	FR662	FR663	FR664	FR665	FR666	FR667	FR668	FR669	FR670	FR671	FR672	FR673	FR674	FR675	FR676	FR677	FR678	FR679	FR680	FR681	FR682	FR683	FR684	FR685	FR686	FR687	FR688	FR689	FR690	FR691	FR692	FR693	FR694	FR695	FR696	FR697	FR698	FR699	FR700	FR701	FR702	FR703	FR704	FR705	FR706	FR707	FR708	FR709	FR710	FR711	FR712	FR713	FR714	FR715	FR716	FR717	FR718	FR719	FR720	FR721	FR722	FR723	FR724	FR725	FR726	FR727	FR728	FR729	FR730	FR731	FR732	FR733	FR734	FR735	FR736	FR737	FR738	FR739	FR740	FR741	FR742	FR743	FR744	FR745	FR746	FR747	FR748	FR749	FR750	FR751	FR752	FR753	FR754	FR755	FR756	FR757	FR758	FR759	FR760	FR761	FR762	FR763	FR764	FR765	FR766	FR767	FR768	FR769	FR770	FR771	FR772	FR773	FR774	FR775	FR776	FR777	FR778	FR779	FR780	FR781	FR782	FR783	FR784	FR785	FR786	FR787	FR788	FR789	FR790	FR791	FR792	FR793	FR794	FR795	FR796	FR797	FR798	FR799	FR800	FR801	FR802	FR803	FR804	FR805	FR806	FR807	FR808	FR809	FR810	FR811	FR812	FR813	FR814	FR815	FR816	FR817	FR818	FR819	FR820	FR821	FR822	FR823	FR824	FR825	FR826	FR827	FR828	FR829	FR830	FR831	FR832	FR833	FR834	FR835	FR836	FR837	FR838	FR839	FR840	FR841	FR842	FR843	FR844	FR845	FR846	FR847	FR848	FR849	FR850	FR851	FR852	FR853	FR854	FR855	FR856	FR857	FR858	FR859	FR860	FR861	FR862	FR863	FR864	FR865	FR866	FR867	FR868	FR869	FR870	FR871	FR872	FR873	FR874	FR875	FR876	FR877	FR878	FR879	FR880	FR881	FR882	FR883	FR884	FR885	FR886	FR887	FR888	FR889	FR890	FR891	FR892	FR893	FR894	FR895	FR896	FR897	FR898	FR899	FR900	FR901	FR902	FR903	FR904	FR905	FR906	FR907	FR908	FR909	FR910	FR911	FR912	FR913	FR914	FR915	FR916	FR917	FR918	FR919	FR920	FR921	FR922	FR923	FR924	FR925	FR926	FR927	FR928	FR929	FR930	FR931	FR932	FR933	FR934	FR935	FR936	FR937	FR938	FR939	FR940	FR941	FR942	FR943	FR944	FR945	FR946	FR947	FR948	FR949	FR950	FR951	FR952	FR953	FR954	FR955	FR956	FR957	FR958	FR959	FR960	FR961	FR962	FR963	FR964	FR965	FR966	FR967	FR968	FR969	FR970	FR971	FR972	FR973	FR974	FR975	FR976	FR977	FR978	FR979	FR980	FR981	FR982	FR983	FR984	FR985	FR986	FR987	FR988	FR989	FR990	FR991	FR992	FR993	FR994	FR995	FR996	FR997	FR998	FR999	FR1000
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54	55	56	57	58	59	60	61	62	63	64	65	66	67	68	69	70	71	72	73	74	75	76	77	78	79	80	81	82	83	84	85	86	87	88	89	90	91	92	93	94	95	96	97	98	99	100	101	102	103	104	105	106	107	108	109	110	111	112	113	114	115	116	117	118	119	120	121	122	123	124	125	126	127	128	129	130	131	132	133	134	135	136	137	138	139	140	141	142	143	144	145	146	147	148	149	150	151	152	153	154	155	156	157	158	159	160	161	162	163	164	165	166	167	168	169	170	171	172	173	174	175	176	177	178	179	180	181	182	183	184	185	186	187	188	189	190	191	192	193	194	195	196	197	198	199	200	201	202	203	204	205	206	207	208	209	210	211	212	213	214	215	216	217	218	219	220	221	222	223	224	225	226	227	228	229	230	231	232	233	234	235	236	237	238	239	240	241	242	243	244	245	246	247	248	249	250	251	252	253	254	255	256	257	258	259	260	261	262	263	264	265	266	267	268	269	270	271	272	273	274	275	276	277	278	279	280	281	282	283	284	285	286	287	288	289	290	291	292	293	294	295	296	297	298	299	300	301	302	303	304	305	306	307	308	309	310	311	312	313	314	315	316	317	318	319	320	321	322	323	324	325	326	327	328	329	330	331	332	333	334	335	336	337	338	339	340	341	342	343	344	345	346	347	348	349	350	351	352	353	354	355	356	357	358	359	360	361	362	363	364	365	366	367	368	369	370	371	372	373	374	375	376	377	378	379	380	381	382	383	384	385	386	387	388	389	390	391	392	393	394	395	396	397	398	399	400	401	402	403	404	405	406	407	408	409	410	411	412	413	414	415	416	417	418	419	420	421	422	423	424	425	426	427	428	429	430	431	432	433	434	435	436	437	438	439	440	441	442	443	444	445	446	447	448																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																								

Figure 11A

19/20

+1 Gln, Val, Gln, Leu, Val, Gln, Ser, Gly, Ala, Glu, Val, Lys, Lys, Pro, Gly, Ser, Ser-
 CAGGTGCAGC TGGTGCAGTC TGGGGCTGAG GTGAAGAAGC CTGGGTCTTC 50
 GTCCACGTCG ACCACGTCAG ACCCCGACTC CACTTCTTCG GACCCAGGAG
 .. Val, Lys, Val, Ser, Cys, Lys, Ala, Ser, Gly, Gly, Thr, Phe, Ser, Ser, His, Ala, Ile-
 GGTGAAGGTC TCCTGCAAGG CTCTCGGAGG CACCTTCAGC AGTCATGCTA 100
 CCACTTCCAG AGGACGTTCC GAAGACCTCC GTGGAAGTCG TCAGTACGAT
 +1 Ser, Trp, Val, Arg, Gln, Ala, Pro, Gly, Gln, Gly, Leu, Glu, Trp, Met, Gly, Asp,
 TCAGCTGGGT GCGACAGGCC CCTGGACAAG GGCTTGAGTG GATGGGAGAC 150
 AGTCGACCCA CGCTGTCCGG GGACCTGTTC CCGAAGTCAC CTACCCCTCG
 +1 Ile, Ile, Pro, Ile, Leu, Gly, Thr, Gly, Asn, Tyr, Ala, Gln, Lys, Phe, Gln, Gly, Arg-
 ATCATCCCTA TCCTTGGTAC AGGAAACTAC GCACAGAAGT TCCAGGGCAG 200
 TAGTAGGGAT AGGAACCATG TCCTTTGATG CGTGTCTTCA AGGTCCCGTC
 +1 Val, Thr, Ile, Thr, Ala, Asp, Glu, Ser, Thr, Ser, Thr, Ala, Tyr, Met, Glu, Leu, Ser
 AGTCACGATT ACCGCGGACG AGTCCACGAG CACAGCCTAC ATGGAGCTGA 250
 TCAGTGCTAA TGGCGCCTGC TCAGGTGCTC GTGTCGGATG TACCTCGACT
 +1 Thr, Leu, Thr, Ser, Glu, Asp, Thr, Ala, Val, Tyr, Tyr, Cys, Glu, Leu, Asp, Trp,
 GCACCCGAC ATCTGAGGAC ACGGCCGTGT ATTACTGTGA ACTTGACTGG 300
 CGTGGGACTG TAGACTCCTG TGCCGGCACA TAATGACACT TGAAGTACCT
 +1 Phe, Tyr, Ile, Trp, Gly, Gln, Gly, Thr, Met, Val, Thr, Val, Ser, Ser,
 TTCTATATCT GGGGCCAAGG GACAATGGTC ACCGTGTCGA GT 350
 AAGATATAGA CCCCAGTTCC CTGTTACCAG TGGCACAGCT CA

Figure 11B

20/20

+1 Glu Val Gln Leu Val Glu Ser Gly Gly Asp Leu Val Gln Pro Gly Gly Ser-
GAGGTGCAGC TGGTGGAGTC TGGGGGAGAC TTGGTACAGC CTGGGGGGTCT 50
CTCCACGTCG ACCACCTCAG ACCCCCTCTG AACCATGTCG GACCCCCCAG

+1 Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Phe Ala Met
CCTGAGACTC TCCTGTGCAG CCTCTGGATT CACCTTTAGC AACTTTGCCA 100
GGACTCTGAG AGGACACGTC GGAGACCTAA GTGGAATCG TTGAAACGGT

+1 Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Trp Val Ala Ala
TGAGCTGGGT CCGCCAGGCT CCCGGGAAGG GGCTGGAGTG GGTCGCGGCT 150
ACTCGACCCA GCGGCTCCGA GGGCCCTTCC CCGACCTCAC CCAGCGCCGA

+1 Ile Gly Gly Arg Ser Gly Thr Thr Phe Tyr Ala Asp Ser Val Lys Gly Arg-
ATTGGCGGTA GAAGTGGTAC CACATTCTAC GCGGACTCCG TGAAGGGCCG 200
TAACCGCCAT CTTACCATG GTGTAAGATG CGCCTGAGGC ACTTCCCGGC

+1 Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Val Tyr Leu Glu Met Asn
GTTCAACCATC TCCAGAGACA ATTCCAAGAA CACGGTCTAT CTGGAATGA 250
CAAGTGGTAG AGGTCTCTGT TAAGGTTCTT GTGCCAGATA GACCTTTACT

+1 Ser Leu Arg Ala Glu Asp Thr Ala Ile Tyr Tyr Cys Ala Lys Arg Gly
ACAGTCTGAG AGCCGAGGAC ACAGCCATTT ATTACTGTGC GAAAAGAGGG 300
TGTCAGACTC TCGGCTCCTG TGTCGGTAAA TAATGACACG CTTTTCTCCC

+1 Arg Gly Gly Tyr Lys Tyr Tyr Gly Met Asp Val Trp Gly Gln Gly Thr Thr-
CGCGGGGGGT ATAAGTATTA TGGGATGGAC GTCTGGGGCC AGGGGACCAC 350
GCGCCCCCA TATTCATAAT ACCCTACCTG CAGACCCCGG TCCCCTGGTG

+1 Val Thr Val Ser Ser
GGTCACCGTG TCGAGT
CCAGTGGCAC AGCTCA 400

Figure 11C

Declaration and Power of Attorney Patent Application (Design or Utility)

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled: "Methods for Diagnosis and Treatment of Haemophilia A Patients with an Inhibitor."

the specification of which

- ☐ is attached hereto
x was filed on November 6, 2000 as application serial no. _____ and or PCT
International Application number PCT/NL99/00285 and was amended on _____ (if
applicable).

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose to the U.S. Patent and Trademark Office all information know to me to be material to patentability as defined in 37 C.F.R. §1.56.

I hereby claim foreign priority benefits under 35 U.S.C. §119(a)-(d) or 35 U.S.C. §365(b) of any foreign application(s) for patent or inventor's certificate, or 35 U.S.C. §365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below any foreign application for patent or inventor's certificate of PCT International application having a filing date before that of the application on which priority is claimed.

Prior Foreign Application(s)		
Number 98201543.0	Country EP	Day/Month/Year Filed 8 May 1998
Number	Country	Day/Month/Year Filed
Number	Country	Day/Month/Year Filed

I hereby claim the benefit under 35 U.S.C. §119(e) of any United States provisional application(s) listed below:

Prior Provisional Application(s)	
Serial Number	Day/Month/Year Filing Date
Serial Number	Day/Month/Year Filing Date
Serial Number	Day/Month/Year Filing Date

I hereby claim the benefit under 35 U.S.C. §120 of any United States application(s), or under 35 U.S.C. §365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. §112, I acknowledge the duty to disclose to the U.S. Patent and Trademark Office all information known to me to be material to patentability as defined in 37 C.F.R. §1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application:

Prior U.S. or International Application(s)		
Serial Number	Day/Month/Year Filed	Status (patented, pending, abandoned)
Serial Number	Day/Month/Year Filed	Status (patented, pending, abandoned)
Serial Number	Day/Month/Year Filed	Status (patented, pending, abandoned)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under 18 U.S.C. §1001 and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Power of Attorney

As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.

Attorney

Registration Number

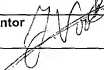
19-
Charles R. Hoffmann 24,102
Ronald J. Baron 29,281
Gerald T. Bodner 30,449
A. Thomas Kammer 28,226
Irving N. Feit 28,601
Alan M. Sack 31,874
Algis Anilionis 36,995
Gregory W. Bachmann 41,593
Anthony E. Bennett 40,910
James F. Harrington P-44,741
Glenn T. Henneberger 36,074
Richard LaCava 41,135
Kevin E. McDermott 35,946
Robert C. Morriss 42,910
Samir R. Patel P-44,998
R. Glenn Schroeder 34,720
Susan A. Sipos 43,128
Roderick S.W. Turner ~~38,639~~
Steven T. Zuschlag 43,309

I hereby authorize them or others whom they may appoint to act and rely on instructions from and communicate directly with the person/organization who/which first sends this case to them and by whom/which I hereby declare that I have consented after full disclosure to be represented unless/until I instructed otherwise.

Please direct all correspondence in this case to at the address indicated below:

Ronald J. Baron
Hoffman & Baron, L.L.P.
6900 Jericho Turnpike
Syosset, New York 11791

1-00

Full Name of Sole or First Inventor		
Family Name <u>Voorberg</u>	First Given Name <u>Johannes</u>	Second Given Name <u>Jacobus</u>
Residence and Citizenship		
City of Residence <u>Assendelft</u>	State or Country of Residence <u>the Netherlands NLX</u>	Country of Citizenship <u>the Netherlands</u>
Post Office Address		
Street Address <u>Klokkemeet 22</u>	City <u>Assendelft</u>	State & Zip Code or Country <u>1566 RH</u>
Signature of Inventor 		Date <u>4/10/2000</u>

2-00

Full Name of Second Inventor, if any		
Family Name <u>van den Brink</u>	First Given Name <u>Edward</u>	Second Given Name <u>Norbert</u>
Residence and Citizenship		
City of Residence <u>Amsterdam</u>	State or Country of Residence <u>the Netherlands NLX</u>	Country of Citizenship <u>the Netherlands</u>
Post Office Address		
Street Address <u>Valkenisseweg 105</u>	City <u>Amsterdam</u>	State & Zip Code or Country <u>1069 BX</u>
Signature of Inventor <u>Eud Brink</u>		Date <u>04/10/00</u>

3-00

Full Name of Third Inventor, if any		
Family Name <u>Turenhout</u>	First Given Name <u>Ellen</u>	Second Given Name <u>Anne Maria</u>
Residence and Citizenship		
City of Residence <u>Noordwijkerhout</u>	State or Country of Residence <u>the Netherlands NLX</u>	Country of Citizenship <u>the Netherlands</u>
Post Office Address		
Street Address <u>Lavendel 19</u>	City <u>Noordwijkerhout</u>	State & Zip Code or Country <u>2211 MV</u>
Signature of Inventor 		Date <u>4/10/2000</u>